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(54) USE OF GENE EXPRESSION MARKERS FOR ASSESSING CARDIAC ALLOGRAFT REJECTION

VERWENDUNG VON GENEXPRESSIONMARKERN ZUR FESTSTELLUNG VON ALLOGRAFT-ABSTOSSUNG DES HERZENS

UTILISATION DE MARQUEURS GENETIQUES D'EXPRESSION POUR L'EVALUATION DU REJET D'UNE ALLOGREFFE CARDIAQUE

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Description

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Related Applications

⁵ **[0001]** This application claims priority to U.S. Application No. 10/131,831, filed April 24, 2002, and U.S. Application No. 10/325,899, filed December 20, 2002.

Field of the Invention

[0002] This invention is in the field of expression profiling following organ transplantation.

Background of the Invention

[0003] Many of the current shortcomings in diagnosis, prognosis, risk stratification and treatment of disease can be approached through the identification of the molecular mechanisms underlying a disease and through the discovery of nucleotide sequences (or sets of nucleotide sequences) whose expression patterns predict the occurrence or progression of disease states, or predict a patient's response to a particular therapeutic intervention. In particular, identification of nucleotide sequences and sets of nucleotide sequences with such predictive value from cells and tissues that are readily accessible would be extremely valuable. For example, peripheral blood is attainable from all patients and can easily be obtained at multiple time points at low cost. This is a desirable contrast to most other cell and tissue types, which are less readily accessible, or accessible only through invasive and aversive procedures. In addition, the various cell types present in circulating blood are ideal for expression profiling experiments as the many cell types in the blood specimen can be easily separated if desired prior to analysis of gene expression. While blood provides a very attractive substrate for the study of diseases using expression profiling techniques, and for the development of diagnostic technologies and the identification of therapeutic targets, the value of expression profiling in blood samples rests on the degree to which changes in gene expression in these cell types are associated with a predisposition to, and pathogenesis and progression of a disease.

[0004] Hematopoiesis is the development and maturation of all cell types of the blood. These include erythrocytes, platelets and leukocytes. Leukocytes are further subdivided into granulocytes (neutrophils, eosinophils, basophils) and mononuclear cells (monocytes, lymphocytes). These cells develop and mature from precursor cells to replenish the circulating pool and to respond to insults and challenges to the system. This occurs in the bone marrow, spleen, thymus, liver, lymph nodes, mucosal associated lymphoid tissue (MALT) and peripheral blood.

[0005] Precursor cells differentiate into immature forms of each lineage and these immature cells develop further into mature cells. This process occurs under the influence and direction of hematopoietic growth factors. When hematopoiesis is stimulated, there is an increase in the number of immature cells in the peripheral blood and in some cases, precursor cells are found at increased frequency. For example, CD34+ cells (hematopoietic stem cells) may increase in frequency in the peripheral blood with an insult to the immune system. For neutrophils, "band" forms are increased, for erythrocytes, reticulocytes or nucleated red cells are seen. Lymphocytes are preceded by lymphoblasts (immature lymphocytes).

[0006] It may be an important clinical goal to measure the rate of production of blood cells of a variety of lineages. Hematological disorders involving over or under production of various blood cells may be treated pharmacologically. For example, anemia (low red blood cells) may be treated with erythropoietin (a hematopoietic growth factor) and response to this therapy can be assessed by measuring RBC production rates. Low neutrophils counts can be treated by administration of G-CSF and this therapy may be monitored by measuring neutrophil production rates. Alternatively, the diagnosis of blood cell disorders is greatly facilitated by determination of lineage specific production rates. For example, anemia (low RBCs) may be caused by decreased cellular production or increased destruction of cells. In the latter case, the rate of cellular production will be increased rather than decreased and the therapeutic implications are very different. Further discussion of the clinical uses of measures of blood cell production rates is given in below.

[0007] Assessment of blood cell production rates may be useful for diagnosis and management of non-hematological disorders. In particular, acute allograft rejection diagnosis and monitoring may benefit from such an approach. Current diagnosis and monitoring of acute allograft rejection is achieved through invasive allograft biopsy and assessment of the biopsy histology. This approach is sub-optimal because of expense of the procedure, cost, pain and discomfort of the patient, the need for trained physician operators, the risk of complications of the procedure, the lack of insight into the functioning of the immune system and variability of pathological assessment. In addition, biopsy can diagnose acute allograft rejection only after significant cellular infiltration into the allograft has occurred. At this point, the process has already caused damage to the allograft. For all these reasons, a simple blood test that can diagnose and monitor acute rejection at an earlier stage in the process is needed. Allograft rejection depends on the presence of functioning cells of the immune system In addition, the process of rejection may cause activation of hematopoiesis. Finally, effective immunosuppressive therapy to treat or prevent acute rejection may suppress hematopoiesis. For these reasons, as-

sessment of hematopoietic cellular production rates may be useful in the diagnosis and monitoring of acute rejection.

[0008] Current techniques for measuring cellular development and production rates are inadequate. The most common approach is to measure the number of mature cells of a lineage of interest over time. For example, if a patient is being treated for anemia (low red blood cell counts), then the physician will order a blood cell count to assess the number of red blood cells (RBCs) in circulation. For this to be effective, the physician must measure the cell count over time and may have to wait 2-4 weeks before being able to assess response to therapy. The same limitation is true for assessment of any cell lineage in the blood.

[0009] An alternative approach is to count the number of immature cells in the peripheral blood by counting them under the microscope. This may allow a more rapid assessment of cellular production rates, but is limited by the need for assessment by a skilled hematologist, observer variability and the inability to distinguish all precursor cells on the basis of morphology alone.

[0010] Bone marrow biopsy is the gold standard for assessment of cellular production rates. In addition to the limitations of the need for skilled physicians, reader variability and the lack of sensitivity of morphology alone, the technique is also limited by the expense, discomfort to the patient and need for a prolonged visit to a medical center. Thus there is a need for a reliable, rapid means for measuring the rate of hematopoeisis in a patient.

[0011] In addition to the relationship between hematopoiesis and variety of disease processes, there is an extensive literature supporting the role of leukocytes, e.g., T-and B-lymphocytes, monocytes and granulocytes, including neutrophils, in a wide range of disease processes, including such broad classes as cardiovascular diseases, inflammatory, autoimmune and rheumatic diseases, infectious diseases, transplant rejection, cancer and malignancy, and endocrine diseases. For example, among cardiovascular diseases, such commonly occurring diseases as atherosclerosis, restenosis, transplant vasculopathy and acute coronary syndromes all demonstrate significant T cell involvement (Smith-Norowitz et al. (1999) Clin Immunol 93:168-175; Jude et al. (1994) Circulation 90:1662-8; Belch et al. (1997) Circulation 95:2027-31). These diseases are now recognized as manifestations of chronic inflammatory disorders resulting from an ongoing response to an injury process in the arterial tree (Ross et al. (1999) Ann Thorac Surg 67:1428-33). Differential expression of lymphocyte, monocyte and neutrophil genes and their products has been demonstrated clearly in the literature. Particularly interesting are examples of differential expression in circulating cells of the immune system that demonstrate specificity for a particular disease, such as arteriosclerosis, as opposed to a generalized association with other inflammatory diseases, or for example, with unstable angina rather than quiescent coronary disease.

[0012] A number of individual genes, e.g., CD11b/CD18 (Kassirer et al. (1999) Am Heart J 138:555-9); leukocyte elastase (Amaro et al. (1995) Eur Heart J 16:615-22; and CD40L (Aukrust et al. (1999) Circulation 100:614-20) demonstrate some degree of sensitivity and specificity as markers of various vascular diseases. In addition, the identification of differentially expressed target and fingerprint genes isolated from purified populations of monocytes manipulated in various in vitro paradigms has been proposed for the diagnosis and monitoring of a range of cardiovascular diseases, see, e.g., US Patents Numbers 6,048,709; 6,087,477; 6,099,823; and 6,124,433 "COMPOSITIONS AND METHODS FOR THE TREATMENT AND DIAGNOSIS OF CARDIOVASCULAR DISEASE" to Falb (see also, WO 97/30065). Lockhart, in US Patent Number 6,033,860 "EXPRESSION PROFILES IN ADULT AND FETAL ORGANS" proposes the use of expression profiles for a subset of identified genes in the identification of tissue samples, and the monitoring of drug effects.

[0013] Morgun et al. (2001) Transplantation Proceedings 1/02(33) identified interferon gamma, TIRC7, perforin and Granzyme B as gene expression markers in blood correlating with rejection of cardiac allografts.

[0014] The accuracy of technologies based on expression profiling for the diagnosis, prognosis, and monitoring of disease would be dramatically increased if numerous differentially expressed nucleotide sequences, each with a measure of specificity for a disease in question, could be identified and assayed in a concerted manner. PCT application WO 02/057414 "LEUKOCYTE EXPRESSION PROFILING" to Wohlgemuth identifies one such set of differentially expressed nucleotides.

[0015] In order to achieve this improved accuracy, the sets of nucleotide sequences once identified need to be validated to identify those differentially expressed nucleotides within a given set that are most useful for diagnosis, prognosis, and monitoring of disease. The present invention addresses these and other needs, and applies to transplant rejection and detection of the rate of hematopoeisis for which differential regulation of genes, or other nucleotide sequences, of peripheral blood can be demonstrated.

Summary of the Invention

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[0016] In order to meet those needs, the present invention is thus directed to a system for detecting differential gene expression.

[0017] In a further variation, the invention is directed to the use of one or more genes to assess cardiac allograft rejection in a patient by detecting the expression level of one or more genes in the patient to diagnose or monitor cardiac transplant rejection in the patient wherein the one or more genes include a nucleotide sequence selected from ID NO:

86, SEQ ID NO:87, SEQ ID NO:107,

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[0018] In another aspect, the methods of diagnosing or monitoring transplant rejection include detecting the expression level of at least two of the genes. In another variation, methods of diagnosing or monitoring transplant rejection include detecting the expression level of at least ten of the genes. In a further variation, the methods of diagnosing or monitoring transplant rejection include detecting the expression level of at least one hundred of the genes. In still a further variation, the methods of diagnosing or monitoring transplant rejection include detecting the expression level of all the listed genes.

[0019] In another aspect, the methods of detecting transplant rejection include detecting the expression level by measuring the RNA level expressed by one or more genes. The method may further including isolating RNA from the patient prior to detecting the RNA level expressed by the one or more genes.

[0020] In one variation, the RNA level is detected by PCR. In a still further variation, the PCR uses primers consisting of nucleotide sequences selected from the group consisting of SEQ ID NO:700, SEQ ID NO:750, SEQ ID NO:751, SEQ ID NO:758, SEQ ID NO:771, SEQ ID NO:1031, SEQ ID NO:1081, SEQ ID NO:1082, SEQ ID NO:1089, SEQ ID NO:1102, SEQ ID NO:1677, SEQ ID NO:1925, SEQ ID NO:1362, SEQ ID NO:1412, SEQ ID NO:1413, SEQ ID NO:1420, SEQ ID NO:1433, SEQ ID NO:2173, The RNA level may be detected by hybridization to the probes. In a further variation, the RNA level is detected by hybridization to an oligonucleotide. Examples of oligonucleotide include oligonucleotides having a nucleotide sequence selected from SEQ ID NO:86, SEQ ID NO:87, SEQ ID NO:107, In still a further variation, the oligonucleotide has the nucleotide sequence SEQ ID NO:94. In another variation, the oligonucleotide has a nucleotide sequence consisting of SEQ ID NO: 107. The oligonucleotide may be DNA, RNA, cDNA, PNA, genomic DNA, or synthetic oligonucleotides.

[0021] In another aspect, the methods of detecting transplant rejection include detecting the expression level by measuring one or more proteins expressed by the one or more genes. In one variation, the one or more proteins include an amino acid sequence selected from SEQ ID NO:2481, SEQ ID NO:2482,

[0022] In another aspect, the method of diagnosing or monitoring cardiac transplant rejection In a patient includes detecting the expression level of one or more genes in the patient to diagnose or monitor cardiac transplant rejection in the patient by measuring one or more proteins expressed by the one or more genes. The one or more proteins may include an amino acid sequence selected from SEQ ID NO:2481, SEQ ID NO:2482, SEQ ID NO:2485, Alternatively, the expression level of the one or more genes may be detected by measuring one or more proteins expressed by one or more genes, and one or more proteins expressed by one or more additional genos. In one variation, the one or more proteins expressed by the one or more genes include an amino acid sequence selected from SEQ ID NO:2481, SEQ ID NO:2482,

[0023] Protein detection may be accomplished by measuring serum. In another variation, the protein is a cell surface protein. In a further variation, the measuring includes using a fluorescent activated cell sorter.

[0024] In another aspect, the disclosure is directed to a substantially purified oligonucleotide having the nucleotide sequence selected from SEQ ID NO:86, SEQ ID NO:87, SEQ ID NO:94, SEQ ID NO:107, and substantially purified oligonucleotides having at least 90% sequence identity to an oligonucleotide having the nucleotide sequence selected from SEQ ID NO:86, SEQ ID NO:87, SEQ ID NO:94, SEQ ID NO:107, In a further aspect, the disclosure is directed to a substantially purified oligonucleotide that hybridizes at high stringency to an oligonucleotide having the nucleotide sequence selected from SEQ ID NO:86, SEQ ID NO:87, SEQ ID NO:94, SEQ ID NO:107, The sequences may be used as diagnostic oligonucleotides for transplant rejection and/or cardiac transplant rejection. The oligonucleotide may have nucleotide sequence including DNA, cDNA, PNA, genomic DNA, or synthetic oligonucleotides.

[0025] In another aspect, the disclosure is directed to a method of diagnosing or monitoring transplant rejection in a patient wherein the expression level of one or more genes in a patient's bodily fluid is detected. In a further variation, the bodily fluid is peripheral blood.

[0026] In another aspect, the disclosure is directed to a method of diagnosing or monitoring transplant rejection in a patient, comprising detecting the expression level of four or more genes in the patient to diagnose or monitor transplant rejection in the patient wherein the four or more genes include a nucleotide sequence selected from SEQ ID NO:86, SEQ ID NO:87, SEQ ID NO:94, SEQ ID NO:107,

[0027] In further the disclosure is also directed to aspect, the disclosure is also directed to a system for detecting gene expression in body fluid including at least two isolated polynucleotides wherein the isolated polynucleotides detect expression of a gene wherein the gene includes a nucleotide sequence selected from SEQ ID NO:86, SEQ ID NO:87, SEQ ID NO:94, SEQ ID NO:107, and the gene in differentially expressed in body fluid in an individual rejecting a transplanted organ compared to the expression of the gene in leukocytes in an individual not rejecting a transplanted organ.

[0028] In another aspect, the disclosure is directed to a system for detecting gene expression in body fluid including at least two isolated polynucleotides wherein the isolated polynucleotides detect expression of a gene wherein the gene includes a nucleotide sequence selected from SEQ ID NO:86, SEQ ID NO:87, SEQ ID NO:94, SEQ ID NO:107, and the gene expression is related to the rate of hematopoiesis or the distribution of hematopoeitic cells along their maturation pathway.

Brief Description of the Figures

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- Figure 1: Figure 1 is a schematic flow chart illustrating a schematic instruction set for characterization of the nucleotide sequence and/or the predicted protein sequence of novel nucleotide sequences.
 - Figure 2: Figure 2 depicts the components of an automated RNA preparation machine.
 - **Figure 3** shows the results of six hybridizations on a mini array graphed (n=6 for each column). The error bars are the SEM. This experiment shows that the average signal from AP prepared RNA is 47% of the average signal from GS prepared RNA for both Cy3 and Cy5.
 - **Figure 4** shows the average background subtracted signal for each of nine leukocyte-specific genes on a mini array. This average is for 3-6 of the above-described hybridizations for each gene. The error bars are the SEM.
 - **Figure 5** shows the ratio of Cy3 to Cy5 signal for a number of genes. After normalization, this ratio corrects for variability among hybridizations and allows comparison between experiments done at different times. The ratio is calculated as the Cy3 background subtracted signal divided by the Cy5 background subtracted signal. Each bar is the average for 3-6 hybridizations. The error bars are SEM. **Figure 6** shows data median Cy3 background subtracted signals for control RNAs using mini arrays. **Figure 7**: Cardiac Allograft rejection diagnostic genes.
 - **A. Example of rejection and no-rejection samples expression data for 5 marker genes**. For each sample, the associated rejection grades are shown as are the expression ratios for 5 differentially expressed genes. The genes are identified by the SEQ ID number for the oligonucleotide. The average fold difference between grade 0 and grade 3A samples is calculated at the bottom.
 - **B. CART classification model**. Decision tree for a 3 gene classification model for diagnosis of cardiac rejection. In the first step, expression of gene 223 is used to divide the patients to 2 branches. The remaining samples in each branch are then further divided by one remaining gene. The samples are classified as either rejection or no rejection. 1 no rejection sample is misclassified as a rejection sample.
 - **C. Surrogates for the CART classification model**. For each of the 3 splitter genes in the CART rejection model described in the example, 5 top surrogate genes are listed that were identified by the CART algorithm.
- 30 Figure 8: Validation of differential expression of a gene discovered using microarrays using real-time PCR
 - **Figure 8A**. The Ct for each patient sample on multiple assays is shown along with the Ct in the R50 control RNA. Triangles represent -RT (reverse transcriptase) controls.
 - **Figure 8B**. The fold difference between the expression of Granzyme B and an Actin reference is shown for 3 samples from patients with and without CMV disease.

Figure 9: Endpoint testing of PCR primers

Electrophoresis and microfluidics are used to assess the product of gene specific PCR primers. β -GUS gel image. Lane 3 is the image for primers F178 and R242. Lanes 2 and 1 correspond to the no-template control and -RT control, respectively.

The electropherogram of β -GUS primers F178 and R242, a graphical representation of Lane 3 from the gel image. β -Actin gel image. Lane 3 is the image for primers F75 and R178. Lanes 2 and 1 correspond to the no-template control and-RT control, respectively.

The electropherogram of β -Actin primers F75 and R178, a graphical representation of Lave 3 from the gel image.

Figure 10: PCR Primer efficiency testing. A standard curve of Ct versus log of the starting RNA amount is shown for 2 genes.

Figure 11: Real-time PCR control gene analysis

11 candidate control genes were tested using real-time PCR on 6 whole blood samples (PAX) paired with 6 mononuclear samples (CPT) from the same patient. Each sample was tested twice. For each gene, the variability of the gene across the samples is shown on the vertical axis (top graph). The average Ct value for each gene is also shown (bottom graph). 2ug RNA was used for PAX samples and 0.5 ug total RNA was used for the mononuclear samples (CPT).

Figure 12: Rejection marker discovery by co-expression with established marker Microarrays were used to measure expression of genes SEQ ID 85 and 302 in samples derived from 240 transplant recipients. For each sample, the expression measurement for 85 is plotted against 302.

Figure 13: ROC (receiver operator characteristics) curve for a 3-gene PCR assay for diagnosis of rejection (see example 17). The Sensitivity and False Positive Rate for each test cutoff is shown.

Brief Description of the Tables

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5 Table 1: Table 1 lists diseases or conditions amenable to study by leukocyte profiling.

Table 2: Transplant Markers

- A. Transplant Genes: Genes useful for monitoring of allograft rejection are listed in this here. The gene symbol and name are given. The NCBI Unigene number (HS) from (Build 160, 16 Feb 2003) is given as is an accession number (ACC) from (Genbank Release 135, 15 April 2003) for an RNA or cDNA is Genbank that corresponds to the gene.
- **B**. Microarray Data: Gene, Gene Name, and ACC are given for each gene as in A (above). Each identified gene has a Non-Parametric Score and Median Rank in NR given from the non-parametric analysis of the data. The genes are ranked from highest to lowest scoring. Down Regulated genes are noted with a 1 in this column.
- **C**. PCR Primers: Primers and probes for real-time PCR assays are given along with their SEQ ID #s. Each gene has 1 or 2 sets of a forward and reverse PCR primer and a hybridization probe for detection in TaqMan or similar assays.
- **D**. PCR Data: Real-time PCR data was generated on a set of transplant samples using sybr green technology as described in the text. For each gene the number of samples (n) used in the analysis is given. An odds ratio and the p-values for a Fisher test and t-test are given for the comparison of acute rejection samples is given (see text).
- **E.** Transplant proteins: For each gene, the corresponding protein in the RefSeq data base (Genbank, Release 135, 18 April 2003) is given (RefSeq Peptide Accession #) along the the SEQ ID for certain proteins for the sequence listing.
- **Table 3**: Viral gene for arrays. Viral genomes were used to design oligonucleotides for the microarrays. The accession numbers for the viral genomes used are given, along with the gene name and location of the region used for oligonucleotide design.
- **Table 4**. Dependent variables for discovery of gene expression markers of cardiac allograft rejection. A stable Grade 0 is a Grade 0 biopsy in a patient who does not experience rejection with the subsequent biopsy. HG or highest grade means that the higher of the biopsy grades from the centralized and local pathologists was used for a definition of the dependent variable.
- **Table 5**: Real-time PCR assay reporter and quencher dyes. Various combinations of reporter and quencher dyes are useful for real-time PCR assays. Reporter and quencher dyes work optimally in specific combinations defined by their spectra. For each reporter, appropriate choices for quencher dyes are given.
- Table 6: Rejection marker PCR assay results
- Results of real-time PCR assays are listed for the comparison of rejection samples to no rejection samples. The fold change is given for expression of each gene in rejection/no rejection samples. The p-value for the t-test comparing the rejection and no rejection classes is given.
- **Table 7**: Summary results of array rejection significance analysis. Summary results are given for correlation analysis of leukocyte gene expression to acute rejection using significance analysis for microarrays (SAM). Five analyses are described. The ISHLT grades used to define the rejection and no rejection classes are given. In each case the highest grade from three pathology reading was taken for analysis. All samples are used for two analyses. The other analyses reduce redundancy of patients used in the analysis by using only one sample per patient ("Non-redundant") or using only one sample per patient within a given class ("Non-redundant within class"). The number of samples used in the analysis is given and the lowest false detection rate (FDR) achieved is noted.
- **Table 9**: Rejection marker sequence analysis. For 63 of the allograft rejection markers listed in Table 2, an analysis of the gene sequence was done. The genes and proteins are identified by accession numbers. The cellular localization of each gene is described as either secreted, nuclear, mitochondrial, cytoplasmic or cellular membrane. The function of the gene is also described.
- **Table 10**: Gene expression markers for immature cells of a variety of lineages are given in Table 10 by way of example **Table 11**: Changes in the rate of hematopoiesis have been correlated to a number of disease states and other pathologies. Examples of such conditions are listed in Table 11.

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Detailed Description of the Invention

Definitions

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[0031] Unless defined otherwise, all scientific and technical terms are understood to have the same meaning as commonly used in the art to which they pertain. For the purpose of the present invention, the following terms are defined below.

[0032] In the context of the invention, the term "gene expression system" refers to any system, device or means to detect gene expression and includes diagnostic agents, candidate libraries, oligonucleotide sets or probe sets.

[0033] The term "monitoring" is used herein to describe the use of gene sets to provide useful information about an individual or an individual's health or disease status. "Monitoring" can include, determination of prognosis, risk-stratification, selection of drug therapy, assessment of ongoing drug therapy, prediction of outcomes, determining response to therapy, diagnosis of a disease or disease complication, following progression of a disease or providing any information relating to a patients health status over time, selecting patients most likely to benefit from experimental therapies with known molecular mechanisms of action, selecting patients most likely to benefit from approved drugs with known molecular mechanisms where that mechanism may be important in a small subset of a disease for which the medication may not have a label, screening a patient population to help decide on a more invasive/expensive test, for example a cascade of tests from a non-invasive blood test to a more invasive option such as biopsy, or testing to assess side effects of drugs used to treat another indication..

[0034] The term "diagnostic oligonucleotide set" generally refers to a set of two or more oligonucleotides that, when evaluated for differential expression of their products, collectively yields predictive data. Such predictive data typically relates to diagnosis, prognosis, monitoring of therapeutic outcomes, and the like. In general, the components of a diagnostic oligonucleotide set are distinguished from nucleotide sequences that are evaluated by analysis of the DNA to directly determine the genotype of an individual as it correlates with a specified trait or phenotype, such as a disease, in that it is the pattern of expression of the components of the diagnostic nucleotide set, rather than mutation or polymorphism of the DNA sequence that provides predictive value. It will be understood that a particular component (or member) of a diagnostic nucleotide set can, in some cases, also present one or more mutations, or polymorphisms that are amenable to direct genotyping by any of a variety of well known analysis methods, e.g., Southern blotting, RFLP, AFLP, SSCP, SNP, and the like.

[0035] A "disease specific target oligonucleotide sequence" is a gene or other oligonucleotide that encodes a polypeptide, most typically a protein, or a subunit of a multi-subunit protein, that is a therapeutic target for a disease, or group of diseases.

[0036] A "candidate library" or a "candidate oligonucleotide library" refers to a collection of oligonucleotide sequences (or gene sequences) that by one or more criteria have an increased probability of being associated with a particular disease or group of diseases. The criteria can be, for example, a differential expression pattern in a disease state or in activated or resting leukocytes in vitro as reported in the scientific or technical literature, tissue specific expression as reported in a sequence database, differential expression in a tissue or cell type of interest, or the like. Typically, a candidate library has at least 2 members or components; more typically, the library has in excess of about 10, or about 100, or about 1000, or even more, members or components.

[0037] The term "disease criterion" is used herein to designate an indicator of a disease, such as a diagnostic factor, a prognostic factor, a factor indicated by a medical or family history, a genetic factor, or a symptom, as well as an overt or confirmed diagnosis of a disease associated with several indicators such as those selected from the above list. A disease criterian includes data describing a patient's health status, including retrospective or prospective health data, e.g. in the form of the patient's medical history, laboratory test results, diagnostic test result, clinical events, medications, lists, response(s) to treatment and risk factors, etc.

[0038] The terms "molecular signature" or "expression profile" refers to the collection of expression values for a plurality (e.g., at least 2, but frequently about 10, about 100, about 1000, or more) of members of a candidate library. In many cases, the molecular signature represents the expression pattern for all of the nucleotide sequences in a library or array of candidate or diagnostic nucleotide sequences or genes. Alternatively, the molecular signature represents the expression pattern for one or more subsets of the candidate library. The term "oligonucleotide" refers to two or more nucleotides. Nucleotides may be DNA or RNA, naturally occurring or synthetic.

[0039] The term "healthy individual," as used herein, is relative to a specified disease or disease criterion. That is, the individual does not exhibit the specified disease criterion or is not diagnosed with the specified disease. It will be understood, that the individual in question, can, of course, exhibit symptoms, or possess various indicator factors for another disease.

[0040] Similarly, an "individual diagnosed with a disease" refers to an individual diagnosed with a specified disease (or disease criterion). Such an individual may, or may not, also exhibit a disease criterion associated with, or be diagnosed with another (related or unrelated) disease.

[0041] An "array" is a spatially or logically organized collection, e.g., of oligonucleotide sequences or nucleotide sequence products such as RNA or proteins encoded by an oligonucleotide sequence. In some embodiments, an array includes antibodies or other binding reagents specific for products of a candidate library.

[0042] When referring to a pattern of expression, a "qualitative" difference in gene expression refers to a difference that is not assigned a relative value. That is, such a difference is designated by an "all or nothing" valuation. Such an all or nothing variation can be, for example, expression above or below a threshold of detection (an on/off pattern of expression). Alternatively, a qualitative difference can refer to expression of different types of expression products, e.g., different alleles (e.g., a mutant or polymorphic allele), variants (including sequence variants as well as post-translationally modified variants), etc.

[0043] In contrast, a "quantitative" difference, when referring to a pattern of gene expression, refers to a difference in expression that can be assigned a value on a graduated scale, (e.g., a 0-5 or 1-10 scale, a + - +++ scale, a grade 1-grade 5 scale, or the like; it will be understood that the numbers selected for illustration are entirely arbitrary and in noway are meant to be interpreted to limit the invention).

15 Gene Expression Systems

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[0044] The disclosure relates to a gene expression system having one or more DNA molecules wherein the one or more DNA molecules has a nucleotide sequence which detects expression of a gene corresponding to the oligonucleotides depicted in the Sequence Listing. In one format, the oligonucleotide detects expression of a gene that is differentially expressed in leukocytes. The gene expression system may be a candidate library, a diagnostic agent, a diagnostic oligonucleotide set or a diagnostic probe set. The DNA molecules may be genomic DNA, protein nucleic acid (PNA), cDNA or synthetic oligonucleotides. Following the procedures taught herein, one can sequences of interest for analyzing gene expression in leukocytes. Such sequences may be predictive of a disease state.

Diagnostic oligonucleotides of the invention

[0045] The disclosure relates to diagnostic nucleotide set(s) comprising members of the leukocyte candidate library listed in Table 2, and in the Sequence Listing, for which a correlation exists between the health status of an individual, the individual's expression of RNA or protein products corresponding to the nucleotide sequence, and the diagnosis and prognosis of transplant rejection. In some instances, only one oligonucleotide is necessary for such detection. Members of a diagnostic oligonucleotide set maybe identified by any means capable of detecting expression of RNA or protein products, including but not limited to differential expression screening, PCR, RT-PCR, SAGE analysis, high-throughput sequencing, microarrays, liquid or other arrays, protein-based methods (e.g., western blotting, proteomics, and other methods described herein), and data mining methods, as further described herein.

[0046] Also described is diagnostic oligonucleotide set comprises at least two oligonucleotide sequences listed in Table 2, or the Sequence Listing which are differentially expressed in leukocytes in an individual with at least one disease criterion for at least one leukocyte-implicated disease relative to the expression in individual without the at least one disease criterion, wherein expression of the two or more nucleotide sequences is correlated with at least one disease criterion, as described below.

[0047] Further disclosed is a diagnostic nucleotide set that comprises at least one oligonucleotide having an oligonucleotide sequence listed in Table 2, or the Sequence Listing which is differentially expressed, and further wherein the differential expression/correlation has not previously been described. In some embodiments, the diagnostic nucleotide set is immobilized on an array.

[0048] In one embodiment, diagnostic nucleotides (or nucleotide sets) are related to the members of the leukocyte candidate library listed in Table 2, or in the Sequence Listing, for which a correlation exists between the health status, diagnosis, and prognosis of transplant rejection (or disease criterion) of an individual. The diagnostic nucleotides are partially or totally contained in (or derived from) full-length gene sequences (or predicted full-length gene sequences) for the members of the candidate library listed inTable 2, and the sequence listing. In some cases, oligonucleotide sequences are designed from EST or Chromosomal sequences from a public database. In these cases the full-length gene sequences may not be known. Full-length sequences in these cases can be predicted using gene prediction algorithms. Alternatively the full-length can be determined by cloning and sequencing the full-length gene or genes that contain the sequence of interest using standard molecular biology approaches described here. The same is true for olignonucleotides designed from our sequencing of cDNA libraries where the cDNA does not match any sequence in the public databases.

[0049] The diagnostic nucleotides may also be derived from other genes that are coexpressed with the correlated sequence or full-length gene. Genes may share expression patterns because they are regulated in the same molecular pathway. Because of the similarity of expression behavior genes are identified as surrogates in that they can substitute for a diagnostic gene in a diagnostic gene set. Example 4 demonstrates the discovery of surrogates from the data and

the sequence listing identifies and gives the sequence for surrogates for cardiac diagnostic genes.

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[0050] As used herein the term "gene cluster" or "cluster" refers to a group of genes related by expression pattern. In other words, a cluster of genes is a group of genes with similar regulation across different conditions, such as graft non-rejection verus graft rejection. The expression profile for each gene in a cluster should be correlated with the expression profile of at least one other gene in that cluster. Correlation may be evaluated using a variety of statistical methods. As used herein the term "surrogate" refers to a gene with an expression profile such that it can substitute for a diagnostic gene in a diagnostic assay. Such genes are often members of the same gene cluster as the diagnostic gene. For each member of a diagnostic gene set, a set of potential surrogates can be identified through identification of genes with similar expression patterns as described below.

[0051] Many statistical analyses produce a correlation coefficient to describe The relatedness between two gene expression patterns. Patterns maybe considered correlated if the correlation coefficient is greater than or equal to 0.8. In preferred embodiments, the correlation coefficient should be greater than 0.85, 0.9 or 0.95. Other statistical methods produce a measure of mutual information to describe the relatedness between two gene expression patterns. Patterns may be considered correlated if the normalized mutual information value is greater than or equal to 0.7. In preferred embodiments, the normalized mutual information value should be greater than 0.8, 0.9 or 0.95, Patterns may also be considered similar if they cluster closely upon hierarchical clustering of gene expression data (Eisen et al. 1998). Similar patterns may be those genes that are among the 1, 2, 5, 10,20,50 or 100 nearest neighbors in a hierarchical clustering or have a similarity score (Eisen ct al. 1998) of > 0.5, 0.7, 0.8, 0.9, 0.95 or 0.99. Similar patterns may also be identified as those genes found to be surrogates in a classification tree by CART (Breiman et al. 1994). Often, but not always, members of a gene cluster have similar biological functions in addition to similar gene expression patterns.

[0052] Correlated genes, clusters and surrogates are identified for the diagnostic genes of the invention. These surrogates may be used as diagnostic genes in an assay instead of, or in addition to, the diagnostic genes for which they are surrogates.

[0053] The disclosure provides diagnostic probe acts. It is understood that a probe includes any reagent capable of specifically identifying a nucleotide sequence of the diagnostic nucleotide set, including but not limited to amplified DNA, amplified RNA, cDNA, synthetic oligonucleotide, partial or full-length nucleic acid sequences. In addition, the probe may identify the protein product of a diagnostic nucleotide sequence, including, for example, antibodies and other affinity reagents.

[0054] It is also understood that each probe can correspond to one gene, or multiple probes can correspond to one gene, or both, or one probe can correspond to more than one gene.

[0055] Homologs and variants of the disclosed nucleic acid molecules may be used in the present invention. Homologs and variants of these nucleic acid molecules will possess a relatively high degree of sequence identity when aligned using standard methods. The sequences encompassed by the invention have at least 40-50, 50-60, 70-80, 80-85, 85-90, 90-95 or 95-100% sequence identity to the sequences disclosed herein.

[0056] It is understood that for expression profiling, variations in the disclosed sequences will still permit detection of gene expression. The degree of sequence identity required to detect gene expression varies depending on the length of the oligomer. For a 60 mer, 6-8 random mutations or 6-8 random deletions in a 60 mer do not affect gene expression detection. Hughes, TR, et al. "Expression profiling using microarrays fabricated by an ink-jet oligonucleotide synthesizer. Nature Biotechnology, 19:343-347(2001). As the length of the DNA sequence is increased, the number of mutations or deletions permitted while still allowing gene expression detection is increased.

[0057] As will be appreciated by those skilled in the art, the sequences of the present invention may contain sequencing errors. That is, there may be incorrect nucleotides, frameshifts, unknown nucleotides, or other types of sequencing errors in any of the sequences; however, the correct sequences will fall within the homology and stringency definitions herein. [0058] The minimum length of an oligonucleotide probe necessary for specific hybridization in the human genome can be estimated using two approaches. The first method uses a statistical argument that the probe will be unique in the human genome by chance. Briefly, the number of independent perfect matches (Po) expected for an oligonucleotide of length L in a genome of complexity C can be calculated from the equation (Laird CD, Chromosoma 32:378 (1971):

Po=(1/4)^L * 2C

[0059] In the case of mammalian genomes, 2C = -3.6 X 10⁹, and an oligonucleotide of 14-15 nucleotides is expected to be represented only once in the genome. However, the distribution of nucleotides in the coding sequence of mammalian genomes is nonrandom (Lathe, R. J. Mol. Biol. 183:1 (1985) and longer oligonucleotides may be preferred in order to in increase the specificity of hybridization. In practical terms, this works out to probes that are 19-40 nucleotides long (Sambrook J et al., infra). The second method for estimating the length of a specific probe is to use a probe long enough to hybridize under the chosen conditions and use a computer to search for that sequence or close matches to the

sequence in the human genome and choose a unique match. Probe sequences are chosen based on the desired hybridization properties as described in Chapter 11 of Sambrook et al, infra. The PRIMER3 program is useful for designing these probes (S. Rozen and H. Skaletsky 1996,1997; Primer3 code available at the web site located at genome.wi.mit.edu/genome_software/other/primer3.html). The sequences of these probes are then compared pair wise against a database of the human genome sequences using a program such as BLAST or MEGABLAST (Madden, T.L et al.(1996) Meth. Enzymol. 266:131-141). Since most of the human genome is now contained in the database, the number of matches will be determined. Probe sequences are chosen that are unique to the desired target sequence.

[0060] In some embodiments, a diagnostic probe set is immobilized on an array. The array is optionally comprises one or more of: a chip array, a plate array, a bead array, a pin array, a membrane array, a solid surface array, a liquid array, an oligonucleotide array, a polynucleotide array or a cDNA array, a microtiter plate, a pin array, a bead array, a membrane or a chip.

[0061] In some embodiments, the leukocyte-implicated disease is selected from the diseases listed in Table 1. In other embodiments, In some embodiments, the disease is atherosclerosis or cardiac allograft rejection. In other embodiments, the disease is congestive heart failure, angina, and myocardial infarction.

[0062] In some embodiments, diagnostic nucleotides of the disclosure are used as a diagnostic gene set in combination with genes that are know to be associated with a disease state ("known markers"). The use of the diagnostic nucleotides in combination with the known markers can provide information that is not obtainable through the known markers alone. The known markers include those identified by the prior art listing provided.

20 <u>Hematopoeisis</u>

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[0063] The present disclosure is also directed to methods of measurement of the rate of hematopoiesis using the diagnostic oligonucleotides of the disclosure and measurement of the rates of hematopoesis by any technique as a method for the monitoring and diagnosis of transplant rejection. Precursor and immature cells often have cell specific phenotypic markers. These are genes and/or proteins that expressed in a restricted manner in immature or precursor cells. This expression decreases with maturation. Gene expression markers for immature cells of a variety of lineages are given in Table 10 below by way of example.

Table 10:

	Table 10:	
Gene	Cell type	
CD10	B-lymphoblasts	
RAG1	B-lymphoblasts	
RAG2	B-lymphoblasts	
NF-E2	Platelets/Megakaryocyte/Erythroid	
GATA-1	Platelets/Megakaryocyte	
GP IIb	Platelets	
pf4	Platelets	
EPO-R	Erythroblast	
Band 4.1	Erythrocyte	
ALAS2	Erythroid specific heme biosynthesis	
hemoglobin chains	Erythocyte	
2,3-BPG mutase	Erythrocyte	
CD16b	Neutrophil	
LAP	Neutrophil	
CD16	NK cells	
CD159a	NK cells	

[0064] By measuring the levels of these and other genes in peripheral blood samples, an assessment of the number and proportion of immature or precursor cells can be made. Of particular use is RNA quantification in erythrocytes and platelets. These cells are anucleated in their mature forms. During

	Anemia - hemolytic	Erythrocyte	Increased	Immunosuppression, Splenectomy
5	Anemia - Renal failure	Erythrocyte	Decreased	Erythropoietin
	Anemia - Chronic disease	Erythrocyte	Decreased	Treat underlying cause
	Polycythemin rubra vera	Erythrocyte	Increased	
10	Idiophic Thrrombocytopenic purpura	Platelet	Increased	Immunosuppression, Splenectomy
15	Thrombotic Thrombocytopenic purpura	Platelet	Increased or decreased	Immunosuppression, plasmapheresis
	Essential thrombocytosis	Platelet	Increased	
	Leukemia	All lineages, variable	Increase, decreased or abnomal	Chemotherapy, BMT
20	Cytopenias due to immunosupression	All lineages, variable	Decreased	Epo,neupogen
	Cytopenias due to Chemotherapy	All lineages, variable	Decreased	Epo, GCSF, GMCSF
25	GVHD	All lineages, variable	Decreased	Immunosuppression
	Myelodysplasia	All lineages, variable	Decreased, increased or abnormal	Chemo?
-	Allograft rejection	Lymphocytes, All lineages	Increased	Immunosuppression
30	Autoimmune diseases (many)	Lymphocytes, All lineages	Increased	Immunosuppression

[0065] The methods described are also useful for monitoring treatment regimens of diseases or other pathologies which are correlated with changes in the rate of hematopoiesis. Furthermore, the methods may be used to monitor treatment with agents that affect the rate of hematopoiesis. One of skill in the art is aware of many such agents. The following agents are examples of such.

[0066] Erythropoietin is a growth factor that is used to treat a variety of anemias that are due to decreased red cell production. Monitoring of red cell production by gene expression or other means may improve dosing and provide a means for earlier assessment of response to therapy for this expensive drug.

[0067] Neupogen (G-CSF) is used for the treatment of low neutrophil counts (neutropenia) usually related to immunosuppression or chemotherapy. Monitoring neutrophil production by gene expression testing or another means may improve dosing, patient selection, and shorten duration of therapy.

[0068] Prednisone I Immunosuppression - One of most common side effects of immunosuppression is suppression of hematopoiesis. This may occur in any cell lineage, Gene expression monitoring or other measures of hematopoietic rates could be used to monitor regularly for cytopenias in a particular cell line and the information could be used to modify dosing, modify therapy or add n specific hematologic growth factor. Following cell counts themselves is less sensitive and results in the need for prolonged trials of therapies at a given dose before efficacy and toxicity can be assessed.

[0069] Monitoring of chemotherapeutic agents -Most chemotherapy agents suppress the bone marrow for some or all lineages. Gene expression testing or other means of assessing hematopoietic rates could be used to monitor regularly for cytopenias in a particular cell line and use information to modify dosing, modify therapy or add a specific hematologic growth factor.

General Molecular Biology References

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[0070] In the context of the invention, nucleic acids and/or proteins are manipulated according to well known molecular biology techniques. Detailed protocols for numerous such procedures are described in, e.g., in Ausubel et al. Current Protocols in Molecular Biology (supplemented through 2000) John Wiley & Sons, New York ("Ausubel"); Sambrook et

al. Molecular Cloning - A Laboratory Manual (2nd Ed.), Vol. 1-3, Cold Spring Harbor Laboratory, Cold Spring Harbor, New York, 1989 ("Sambrook"), and Berger and Kimmel Guide to Molecular Cloning Techniques, Methods in Enzymology volume 152 Academic Press, Inc., San Diego, CA ("Berger").

[0071] In addition to the above references, protocols for in vitro amplification techniques, such as the polymerase chain reaction (PCR), the ligase chain reaction (LCR), Q-replicase amplification, and other RNA polymerase mediated techniques (e.g., NASBA), useful e.g., for amplifying cDNA probes of the invention, are found in Mullis et al. (1987) U.S. Patent No. 4,683,202; PCR Protocols A Guide to Methods and Applications (Innis et al. eds) Academic Press Inc. San Diego, CA (1990) ("Innis"); Arnheim and Levinson (1990) C&EN 36; The Journal Of NIH Research (1991) 3:81; Kwoh et al. (1989) Proc Natl Acad Sci USA 86, 1173; Guatelli et al. (1990) Proc Natl Acad Sci USA 87:1874; Lomell et al. (1989) J Clin Chem 35:1826; Landegren et al. (1988) Science 241:1077; Van Brunt (1990) Biotechnology 8:291; Wu and Wallace (1989) Gene 4: 560; Barringer et al. (1990) Gene 89:117, and Sooknanan and Malek (1995) Biotechnology 13:563. Additional methods, useful for cloning nucleic acids in the context of the present invention, include Wallace et al. U.S. Pat. No. 5,426,039. Improved methods of amplifying large nucleic acids by PCR are summarized in Cheng et al. (1994) Nature 369:684 and the references therein.

[0072] Certain polynucleotides of the invention, e.g., oligonucleotides can be synthesized utilizing various solid-phase strategies involving mononucleotide- and/or trinucleotide-based phosphoramidite coupling chemistry. For example, nucleic acid sequences can be synthesized by the sequential addition of activated monomers and/or trimers to an elongating polynucleotide chain. See e.g., Caruthers, M.H. et al. (1992) Meth Enzymol 211:3.

[0073] In lieu of synthesizing the desired sequences, essentially any nucleic acid can be custom ordered from any of a variety of commercial sources, such as The Midland Certified Reagent Company, The Great American Gene Company ExpressGen, Inc., Operon Technologies, Inc. and many others.

[0074] Similarly, commercial sources for nucleic acid and protein microarrays are available, and include, e.g., Agilent Technologies, Palo Alto, CA Affymetrix, Santa Clara, CA; and others.

[0075] One area of relevance to the present invention is hybridization of oligonucleotides. Those of skill in the art differentiate hybridization conditions based upon the stringency of hybridization. For example, highly stringent conditions could include hybridization to filter-bound DNA in 0.5 M NaHPO₄, 7% sodium dodecyl sulfate (SDS), 1 mM EDTA at 65° C, and washing in 0.1XSSC/0.1% SDS at 68° C. (Ausubel F. M. et al., eds., 1989, Current Protocols in Molecular Biology, Vol. I, Green Publishing Associates, Inc., and John Wiley & sons, Inc., New York, at p. 2.10.3). Moderate stringency conditions could include, e.g., washing in 0.2XSSC/0.1 % SDS at 42°C. (Ausubel et al., 1989, supra).

The disclosure also includes nucleic acid molecules, preferably DNA molecules, that hybridize to, and are therefore the complements of, the DNA sequences of the present disclosure. Such hybridization conditions may be highly stringent or less highly stringent, as described above. In instances wherein the nucleic acid molecules are deoxyoligonucleotides ("oligos"), highly stringent conditions may refer, e.g., to washing in 6xSSC/0.05% sodium pyrophosphate at 37°C. (for 14-base oligos), 48°C. (for 17-base oligos), 55°C. (for 20-base oligos), and 60°C. (for 23-base oligos). These nucleic acid molecules may act as target nucleotide sequence antisense molecules, useful, for example, in target nucleotide sequence regulation and/or as antisense primers in amplification reactions of target nucleotide sequence nucleic acid sequences. Further, such sequences may be used as part of ribozyme and/or triple helix sequences, also useful for target nucleotide sequence regulation. Still further, such molecules may be used as components of diagnostic methods whereby the presence of a disease-causing allele, may be detected.

Identification of diagnostic nucleotide sets

Candidate library

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[0076] Libraries of candidates that are differentially expressed in leukocytes are substrates for the identification and evaluation of diagnostic oligonucleotide sets and disease specific target nucleotide sequences.

[0077] The term leukocyte is used generically to refer to any nucleated blood cell that is not a nucleated erythrocyte. More specifically, leukocytes can be subdivided into two broad classes. The first class includes granulocytes, including, most prevalently, neutrophils, as well as eosinophils and basophils at low frequency. The second class, the non-granular or mononuclear leukocytes, includes monocytes and lymphocytes (e.g., T cells and B cells). There is an extensive literature in the art implicating leukocytes, e.g., neutrophils, monocytes and lymphocytes in a wide variety of disease processes, including inflammatory and rheumatic diseases, neurodegenerative diseases (such as Alzheimer's dementia), cardiovascular disease, endocrine diseases, transplant rejection, malignancy and infectious diseases, and other diseases listed in Table 1. Mononuclear cells are involved in the chronic immune response, while granulocytes, which make up approximately 60% of the leukocytes, have a non-specific and stereotyped response to acute inflammatory stimuli and often have a life span of only 24 hours.

[0078] In addition to their widespread involvement and/or implication in numerous disease related processes, leukocytes are particularly attractive substrates for clinical and experimental evaluation for a variety of reasons. Most impor-

tantly, they are readily accessible at low cost from essentially every potential subject. Collection is minimally invasive and associated with little pain, disability or recovery time. Collection can be performed by minimally trained personnel (e.g., phlebotomists, medical technicians, etc.), in a variety of clinical and non-clinical settings without significant technological expenditure. Additionally, leukocytes are renewable, and thus available at multiple time points for a single subject.

Assembly of an initial candidature library

[0079] The initial candidate library was assembled from a combination of "mining" publication and sequence databases and construction of a differential expression library. Candidate oligonucleotide sequences in the library may be represented by a full-length or partial nucleic acid sequence, deoxyribonucleic acid (DNA) sequence, cDNA sequence, RNA sequence, synthetic oligonucleotides, etc. The nucleic acid sequence can be at least 19 nucleotides in length, at least 25 nucleotides, at least 40 nucleotides, at least 100 nucleotides, or larger. Alternatively, the protein product of a candidate nucleotide sequence may be represented in a candidate library using standard methods, as further described below. In selecting and validatating diagnostic oligonucleotides, an initial library of 8,031 candidate oligonucleotide sequences using nucleic acid sequences of 50 nucleotides in length was constructed as described below.

Candidate nucleotide library

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[0080] We identified members of an initial candidate nucleotide library that are differentially expressed in activated leukocytes and resting leukocytes. From that initial candidate nucleotide library, a pool of candidates was selected as listed in Table 2, and the sequence listing. Accordingly, the disclosure provides the candidate leukocyte nucleotide library comprising the nucleotide sequences listed in Table 2, and in the sequence listing. In another embodiment, the disclosure provides an candidate library comprising at least one nucleotide sequence listed in Table 2 and the sequence listing. In another embodiment, the disclosure provides an candidate library comprising at least two nucleotide sequences listed in Table 2 and the sequence listing. In another embodiment, the at least two nucleotide sequence are at least 19 nucleotides in length, at least 35 nucleotides, at least 40 nucleotides or at least 100 nucleotides. In some embodiments, the nucleotide sequences comprises deoxyribonucleic acid (DNA) sequence, ribonucleic acid (RNA) sequence, synthetic oligonucleotide sequence, or genomic DNA sequence. It is understood that the nucleotide sequences may each correspond to one gene, or that several nucleotide sequences may correspond to one gene, or both.

[0081] The disclosure also provides probes to the candidate nucleotide library. The probes can comprise at least two nucleotide sequences listed in Table 2, or the sequence listing which are differentially expressed in leukocytes in an individual with a least one disease criterion for at least one leukocyte-related disease and in leukocytes in an individual without the at least one disease criterion, wherein expression of the two or more nucleotide sequences is correlated with at least one disease criterion. It is understood that a probe may detect either the RNA expression or protein product expression of the candidate nucleotide library. Alternatively, or in addition, a probe can detect a genotype associated with a candidate nucleotide sequence, as further described below. The probes for the candidate nucleotide library can also be immobilized on an array.

[0082] The candidate nucleotide library disclosed is useful in identifying diagnostic nucleotide sets of the invention. The candidate nucleotide sequences may be further characterized, and may, be identified as a disease target nucleotide sequence and/or a novel nucleotide sequence, as described below. The candidate nucleotide sequences may also be suitable for use as imaging reagents, as described below.

Detection of non-leukocyte expressed genes

[0083] When measuring gene expression levels in a blood sample, RNAs may be measured that are not derived from leukocytes. Examples are viral genes, free RNAs that have been released from damaged non-leukocyte cell types or RNA from circulating non-leukocyte cell types. For example, in the process of acute allograft rejection, tissue damage may result in release of allograft cells or RNAs derived from allograft cells into the circulation. In the case of cardiac allografts, such transcripts may be specific to muscle (myoglobin) or to cardiac muscle (Troponin I, Toponin T, CK-MB). Presence of cardiac specific mRNAs in peripheral blood may indicate ongoing or recent cardiac cellular damage (resulting from acute rejection). Therefore, such genes may be excellent diagnostic markers for allograft rejection.

Generation of Expression Patterns

RNA, DNA or protein sample procurement

[0084] Following identification or assembly of a library of differentially expressed candidate nucleotide sequences,

leukocyte expression profiles corresponding to multiple members of the candidate library are obtained. Leukocyte samples from one or more subjects are obtained by standard methods. Most typically, these methods involve trans-cutaneous venous sampling of peripheral blood. While sampling of circulating leukocytes from whole blood from the peripheral vasculature is generally the simplest, least invasive, and lowest cost alternative, it will be appreciated that numerous alternative sampling procedures exist, and are favorably employed in some circumstances. No pertinent distinction exists, in fact, between leukocytes sampled from the peripheral vasculature, and those obtained, e.g., from a central line, from a central artery, or indeed from a cardiac catheter, or during a surgical procedure which accesses the central vasculature. In addition, other body fluids and tissues that are, at least in part, composed of leukocytes are also desirable leukocyte samples. For example, fluid samples obtained from the lung during bronchoscopy may be rich in leukocytes, and amenable to expression profiling in the context of the invention, e.g., for the diagnosis, prognosis, or monitoring of lung transplant rejection, inflammatory lung diseases or infectious lung disease. Fluid samples from other tissues, e.g., obtained by endoscopy of the colon, sinuses, esophagus, stomach, small bowel, pancreatic duct, biliary tree, bladder, ureter, vagina, cervix or uterus, etc., are also suitable. Samples may also be obtained other sources containing leukocytes, e.g., from urine, bile, cerebrospinal fluid, feces, gastric or intestinal secretions, semen, or solid organ or joint biopsies. [0085] Most frequently, mixed populations of leukocytes, such as are found in whole blood are utilized in the methods of the present invention. A crude separation, e.g., of mixed leukocytes from red blood cells, and/or concentration, e.g., over a sucrose, percoll or ficoll gradient, or by other methods known in the art, can be employed to facilitate the recovery of RNA or protein expression products at sufficient concentrations, and to reduce non-specific background. In some instances, it can be desirable to purify sub-populations of leukocytes, and methods for doing so, such as density or affinity gradients, flow cytometry, fluorescence Activated Cell Sorting (FACS), immuno-magnetic separation, "panning," and the like, are described in the available literature and below.

Obtaining DNA, RNA and protein samples for expression profiling

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[0086] Expression patterns can be evaluated at the level of DNA, or RNA or protein products. For example, a variety of techniques are available for the isolation of RNA from whole blood. Any technique that allows isolation of mRNA from cells (in the presence or absence of rRNA and tRNA) can be utilized. In brief, one method that allows reliable isolation of total RNA suitable for subsequent gene expression analysis, is described as follows. Peripheral blood (either venous or arterial) is drawn from a subject, into one or more sterile, endotoxin free, tubes containing an anticoagulant (e.g., EDTA, citrate, heparin, etc.). Typically, the sample is divided into at least two portions. One portion, e.g., of 5-8 ml of whole blood is frozen and stored for future analysis, e.g., of DNA or protein. A second portion, e.g., of approximately 8 ml whole blood is processed for isolation of total RNA by any of a variety of techniques as described in, e.g, Sambook, Ausubel, below, as well as U.S. Patent Numbers: 5,728,822 and 4,843,155.

[0087] Typically, a subject sample of mononuclear leukocytes obtained from about 8 ml of whole blood, a quantity readily available from an adult human subject under most circumstances, yields 5-20 µg of total RNA. This amount is ample, e.g., for labeling and hybridization to at least two probe arrays. Labeled probes for analysis of expression patterns of nucleotides of the candidate libraries are prepared from the subject's sample of RNA using standard methods. In many cases, cDNA is synthesized from total RNA using a polyT primer and labeled, e.g., radioactive or fluorescent, nucleotides. The resulting labeled cDNA is then hybridized to probes corresponding to members of the candidate nucleotide library, and expression data is obtained for each nucleotide sequence in the library. RNA isolated from subject samples (e.g., peripheral blood leukocytes, or leukocytes obtained from other biological fluids and samples) is next used for analysis of expression patterns of nucleotides of the candidate libraries.

[0088] In some cases, however, the amount of RNA that is extracted from the leukocyte sample is limiting, and amplification of the RNA is desirable. Amplification may be accomplished by increasing the efficiency of probe labeling, or by amplifying the RNA sample prior to labeling. It is appreciated that care must be taken to select an amplification procedure that does not introduce any bias (with respect to gene expression levels) during the amplification process.

[0089] Several methods are available that increase the signal from limiting amounts of RNA, e.g. use of the Clontech (Glass Fluorescent Labeling Kit) or Stratagene (Fairplay Microarray Labeling Kit), or the Micromax kit (New England Nuclear, Inc.). Alternatively, cDNA is synthesized from RNA using a T7- polyT primer, in the absence of label, and DNA dendrimers from Genisphere (3DNA Submicro) are hybridized to the poly T sequence on the primer, or to a different "capture sequence" which is complementary to a fluorescently labeled sequence. Each 3DNA molecule has 250 fluorescent molecules and therefore can strongly label each cDNA.

[0090] Alternatively, the RNA sample is amplified prior to labeling. For example, linear amplification may be performed, as described in U.S. Patent No. 6,132,997. A T7-polyT primer is used to generate the cDNA copy of the RNA. A second DNA strand is then made to complete the substrate for amplification. The T7 promoter incorporated into the primer is used by a T7 polymerase to produce numerous antisense copies of the original RNA. Fluorescent dye labeled nucleotides are directly incorporated into the RNA. Alternatively, amino allyl labeled nucleotides are incorporated into the RNA, and then fluorescent dyes are chemically coupled to the amino allyl groups, as described in Hughes. Other exemplary

methods for amplification are described below.

[0091] It is appreciated that the RNA isolated must contain RNA derived from leukocytes, but may also contain RNA from other cell types to a variable degree. Additionally, the isolated RNA may come from subsets of leukocytes, e.g. monocytes and/or T-lymphocytes, as described above. Such consideration of cell type used for the derivation of RNA depend on the method of expression profiling used. Subsets of leukocytes can be obtained by fluorescence activated cell sorting (FACS), microfluidics cell seperation systems or a variety of other methods. Cell sorting may be necessary for the discovery of diagnostic gene sets, for the implementation of gene sets as products or both. Cell sorting can be achieved with a variety of technologies (See Galbraith et al. 1999, Cantor et al. 1975, see also the technology of Guava Technologies, Hayward, CA).

[0092] DNA samples may be obtained for analysis of the presence of DNA mutations, single nucleotide polymorphisms (SNPs), or other polymorphisms. DNA is isolated using standard techniques, e.g. *Maniatus, supra.*

[0093] Expression of products of candidate nucleotides may also be assessed using proteomics. Protein(s) are detected in samples of patient serum or from leukocyte cellular protein. Serum is prepared by centrifugation of whole blood, using standard methods. Proteins present in the serum may have been produced from any of a variety of leukocytes and non-leukocyte cells, and include secreted proteins from leukocytes. Alternatively, leukocytes or a desired sub-population of leukocytes are prepared as described above. Cellular protein is prepared from leukocyte samples using methods well known in the art, e.g., Trizol (Invitrogen Life Technologies, cat # 15596108; Chomczynski, P. and Sacchi, N. (1987) Anal. Biochem. 162, 156; Simms, D., Cizdziel, P.E., and Chomczynski, P. (1993) Focus® 15, 99; Chomczynski, P., Bowers-Finn, R., and Sabatini, L. (1987) J. of NIH Res. 6, 83; Chomczynski, P. (1993) Bio/Techniques 15, 532; Bracete, A.M., Fox, D.K., and Simms, D. (1998) Focus 20, 82; Sewall, A. and McRae, S. (1998) Focus 20, 36; Anal Biochem 1984 Apr;138(1):141-3, A method for the quantitative recovery of protein in dilute solution in the presence of detergents and lipids; Wessel D, Flugge UI. (1984) Anal Biochem. 1984 Apr;138(1):141-143.

[0094] The assay itself may be a cell sorting assay in which cells are sorted and/or counted based on cell surface expression of a protein marker. (See Cantor et al. 1975, Galbraith et al. 1999)

Obtaining expression patterns

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[0095] Expression patterns, or profiles, of a plurality of nucleotides corresponding to members of the candidate library are then evaluated in one or more samples of leukocytes. Typically, the leukocytes are derived from patient peripheral blood samples, although, as indicated above, many other sample sources are also suitable. These expression patterns constitute a set of relative or absolute expression values for a some number of RNAs or protein products corresponding to the plurality of nucleotide sequences evaluated, which is referred to herein as the subject's "expression profile" for those nucleotide sequences. While expression patterns for as few as one independent member of the candidate library can be obtained, it is generally preferable to obtain expression patterns corresponding to a larger number of nucleotide sequences, e.g., about 2, about 5, about 10, about 20, about 100, about 200, about 500, or about 1000, or more. The expression pattern for each differentially expressed component member of the library provides a finite specificity and sensitivity with respect to predictive value, e.g., for diagnosis, prognosis, monitoring, and the like.

Clinical Studies, Data and Patient Groups

[0096] For the purpose of discussion, the term subject, or subject sample of leukocytes, refers to an individual regardless of health and/or disease status. A subject can be a patient, a study participant, a control subject, a screening subject, or any other class of individual from whom a leukocyte sample is obtained and assessed in the context of the invention. Accordingly, a subject can be diagnosed with a disease, can present with one or more symptom of a disease, or a predisposing factor, such as a family (genetic) or medical history (medical) factor, for a disease, or the like. Alternatively, a subject can be healthy with respect to any of the aforementioned factors or criteria. It will be appreciated that the term "healthy" as used herein, is relative to a specified disease, or disease factor, or disease criterion, as the term "healthy" cannot be defined to correspond to any absolute evaluation or status. Thus, an individual defined as healthy with reference to any specified disease or disease criterion, can in fact be diagnosed with any other one or more disease, or exhibit any other one or more disease criterion.

[0097] Furthermore, while the discussion of the invention focuses, and is exemplified using human sequences and samples, the invention is equally applicable, through construction or selection of appropriate candidate libraries, to non-human animals, such as laboratory animals, e.g., mice, rats, guinea pigs, rabbits; domesticated livestock, e.g., cows, horses, goats, sheep, chicken, etc.; and companion animals, e.g., dogs, cats, etc.

Methods for obtaining expression data

[0098] Numerous methods for obtaining expression data are known, and any one or more of these techniques, singly

or in combination, are suitable for determining expression profiles in the context of the present invention. For example, expression patterns can be evaluated by northern analysis, PCR, RT-PCR, Taq Man analysis, FRET detection, monitoring one or more molecular beacon, hybridization to an oligonucleotide array, hybridization to a cDNA array, hybridization to a polynucleotide array, hybridization to a liquid microarray, hybridization to a microelectric array, molecular beacons, cDNA sequencing, clone hybridization, cDNA fragment fingerprinting, serial analysis of gene expression (SAGE), subtractive hybridization, differential display and/or differential screening (see, e.g., Lockhart and Winzeler (2000) Nature 405:827-836, and references cited therein).

[0099] For example, specific PCR primers are designed to a member(s) of an candidate nucleotide library. cDNA is prepared from subject sample RNA by reverse transcription from a poly-dT oligonucleotide primer, and subjected to PCR. Double stranded cDNA may be prepared using primers suitable for reverse transcription of the PCR product, followed by amplification of the cDNA using in vitro transcription. The product of in vitro transcription is a sense-RNA corresponding to the original member(s) of the candidate library. PCR product may be also be evaluated in a number of ways known in the art, including real-time assessment using detection of labeled primers, e.g. TaqMan or molecular beacon probes. Technology platforms suitable for analysis of PCR products include the ABI 7700, 5700, or 7000 Sequence Detection Systems (Applied Biosystems, Foster City, CA), the MJ Research Opticon (MJ Research, Waltham, MA), the Roche Light Cycler (Roche Diagnositics, Indianapolis, IN), the Stratagene MX4000 (Stratagene, La Jolla, CA), and the Bio-Rad iCycler (Bio-Rad Laboratories, Hercules, CA). Alternatively, molecular beacons are used to detect presence of a nucleic acid sequence in an unamplified RNA or cDNA sample, or following amplification of the sequence using any method, e.g. IVT (In Vitro transcription) or NASBA (nucleic acid sequence based amplification). Molecular beacons are designed with sequences complementary to member(s) of an candidate nucleotide library, and are linked to fluorescent labels. Each probe has a different fluorescent label with non-overlapping emission wavelengths. For example, expression often genes may be assessed using ten different sequence-specific molecular beacons.

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[0100] Alternatively, or in addition, molecular beacons are used to assess expression of multiple nucleotide sequences at once. Molecular beacons with sequence complimentary to the members of a diagnostic nucleotide set are designed and linked to fluorescent labels. Each fluorescent label used must have a non-overlapping emission wavelength. For example, 10 nucleotide sequences can be assessed by hybridizing 10 sequence specific molecular beacons (each labeled with a different fluorescent molecule) to an amplified or un-amplified RNA or cDNA sample. Such an assay bypasses the need for sample labeling procedures.

[0101] Alternatively, or in addition bead arrays can be used to assess expression of multiple sequences at once. See, e.g, LabMAP 100, Luminex Corp, Austin, Texas). Alternatively, or in addition electric arrays are used to assess expression of multiple sequences, as exemplified by the e-Sensor technology of Motorola (Chicago, III.) or Nanochip technology of Nanogen (San Diego, CA.)

[0102] Of course, the particular method elected will be dependent on such factors as quantity of RNA recovered, practitioner preference, available reagents and equipment, detectors, and the like. Typically, however, the elected method(s) will be appropriate for processing the number of samples and probes of interest. Methods for high-throughput expression analysis are discussed below.

[0103] Alternatively, expression at the level of protein products of gene expression is performed. For example, protein expression, in a sample of leukocytes, can be evaluated by one or more method selected from among; western analysis, two-dimensional gel analysis, chromatographic separation, mass spectrometric detection, protein-fusion reporter constructs, colorimetric assays, binding to a protein array and characterization of polysomal mRNA. One particularly favorable approach involves binding of labeled protein expression products to an array of antibodies specific for members of the candidate library. Methods for producing and evaluating antibodies are widespread in the art, see, e.g., Coligan, supra; and Harlow and Lane (1989) Antibodies: A Laboratory Manual, Cold Spring Harbor Press, NY ("Harlow and Lane"). Additional details regarding a variety of immunological and immunoassay procedures adaptable to the present invention by selection of antibody reagents specific for the products of candidate nucleotide sequences can be found in, e.g., Stites and Terr (eds.)(1991) Basic and Clinical Immunology, 7th ed., and Paul, supra. Another approach uses systems for performing desorption spectrometry. Commercially available systems, e.g., from Ciphergen Biosystems, Inc. (Fremont, CA) are particularly well suited to quantitative analysis of protein expression. Indeed, Protein Chip® arrays (see, e.g., the web site ciphergen.com) used in desorption spectrometry approaches provide arrays for defection of protein expression. Alternatively, affinity reagents, e.g., antibodies, small molecules, etc.) are developed that recognize epitopes of the protein product. Affinity assays are used in protein array assays, e.g. to detect the presence or absence of particular proteins. Alternatively, affinity reagents are used to detect expression using the methods described above. In the case of a protein that is expressed on the cell surface of leukocytes, labeled affinity reagents are bound to populations of leukocytes, and leukocytes expressing the protein are identified and counted using fluorescent activated cell sorting (FACS).

[0104] It is appreciated that the methods of expression evaluation discussed herein, although discussed in the context of discovery of diagnostic nucleotide sets, are equally applicable for expression evaluation when using diagnostic nucleotide sets for, e.g. diagnosis of diseases, as further discussed below.

High Throughout Expression Assays

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[0105] A number of suitable high throughput formats exist for evaluating gene expression. Typically, the term high throughput refers to a format that performs at least about 100 assays, or at least about 500 assays, or at least about 1000 assays, or at least about 5000 assays, or at least about 10,000 assays, or more per day. When enumerating assays, either the number of samples or the number of candidate nucleotide sequences evaluated can be considered. For example, a northern analysis of, e.g., about 100 samples performed in a gridded array, e.g., a dot blot, using a single probe corresponding to an candidate nucleotide sequence can be considered a high throughput assay. More typically, however, such an assay is performed as a series of duplicate blots, each evaluated with a distinct probe corresponding to a different member of the candidate library. Alternatively, methods that simultaneously evaluate expression of about 100 or more candidate nucleotide sequences in one or more samples, or in multiple samples, are considered high throughput.

[0106] Numerous technological platforms for performing high throughput expression analysis are known. Generally, such methods involve a logical or physical array of either the subject samples, or the candidate library, or both. Common array formats include both liquid and solid phase arrays. For example, assays employing liquid phase arrays, e.g., for hybridization of nucleic acids, binding of antibodies or other receptors to ligand, etc., can be performed in multiwell, or microtiter, plates. Microtiter plates with 96, 384 or 1536 wells are widely available, and even higher numbers of wells, e.g., 3456 and 9600 can be used. In general, the choice of microtiter plates is determined by the methods and equipment, e.g., robotic handling and loading systems, used for sample preparation and analysis. Exemplary systems include, e.g., the ORCATM system from Beckman-Coulter, Inc. (Fullerton, CA) and the Zymate systems from Zymark Corporation (Hopkinton, MA).

[0107] Alternatively, a variety of solid phase arrays can favorably be employed in to determine expression patterns in the context of the invention. Exemplary formats include membrane or filter arrays (e.g., nitrocellulose, nylon), pin arrays, and bead arrays (e.g., in a liquid "slurry"). Typically, probes corresponding to nucleic acid or protein reagents that specifically interact with (e.g., hybridize to or bind to) an expression product corresponding to a member of the candidate library are immobilized, for example by direct or indirect cross-linking, to the solid support. Essentially any solid support capable of withstanding the reagents and conditions necessary for performing the particular expression assay can be utilized. For example, functionalized glass, silicon, silicon dioxide, modified silicon, any of a variety of polymers, such as (poly)tetrafluoroethylene, (poly)vinylidenedifluoride, polystyrene, polycarbonate, or combinations thereof can all serve as the substrate for a solid phase array.

[0108] In a preferred embodiment, the array is a "chip" composed, e.g., of one of the above specified materials. Polynucleotide probes, e.g., RNA or DNA, such as cDNA, synthetic oligonucleotides, and the like, or binding proteins such as antibodies, that specifically interact with expression products of individual components of the candidate library are affixed to the chip in a logically ordered manner, i.e., in an array. In addition, any molecule with a specific affinity for either the sense or anti-sense sequence of the marker nucleotide sequence (depending on the design of the sample labeling), can be fixed to the array surface without loss of specific affinity for the marker and can be obtained and produced for array production, for example, proteins that specifically recognize the specific nucleic acid sequence of the marker, ribozymes, peptide nucleic acids (PNA), or other chemicals or molecules with specific affinity.

[0109] Detailed discussion of methods for linking nucleic acids and proteins to a chip substrate, are found in, e.g., US Patent No. 5,143,854 "LARGE SCALE PHOTOLITHOGRAPHIC SOLID PHASE SYNTHESIS OF POLYPEPTIDES AND RECEPTOR BINDING SCREENING THEREOF" to Pirrung et al., issued, September 1, 1992; US Patent No. 5,837,832 "ARRAYS OF NUCLEIC ACID PROBES ON BIOLOGICAL CHIPS" to Chee et al., issued November 17, 1998; US Patent No. 6,087,112 "ARRAYS WITH MODIFIED OLIGONUCLEOTIDE AND POLYNUCLEOTIDE COMPOSITIONS" to Dale, issued July 11, 2000; US Patent No. 5,215,882 "METHOD OF IMMOBILIZING NUCLEIC ACID ON A SOLID SUBSTRATE FOR USE IN NUCLEIC ACID HYBRIDIZATION ASSAYS" to Bahl et al., issued June 1, 1993; US Patent No. 5,707,807 "MOLECULAR INDEXING FOR EXPRESSED GENE ANALYSIS" to Kato, issued January 13, 1998; US Patent No. 5,807,522 "METHODS FOR FABRICATING MICROARRAYS OF BIOLOGICAL SAMPLES" to Brown et al., issued September 15, 1998; US Patent No. 5,958,342 "JET DROPLET DEVICE" to Gamble et al., issued Sept. 28, 1999; US Patent 5,994,076 "METHODS OF ASSAYING DIFFERENTIAL EXPRESSION" to Chenchik et al., issued Nov. 30, 1999; US Patent No. 6,004,755 "QUANTITATIVE MICROARRAY HYBRIDIZATION ASSAYS" to Wang, issued Dec. 21, 1999; US Patent No. 6,048,695 "CHEMICALLY MODIFIED NUCLEIC ACIDS AND METHOD FOR COUPLING NUCLEIC ACIDS TO SOLID SUPPORT" to Bradley et al., issued April 11, 2000; US Patent No. 6,060,240 "METHODS FOR MEASURING RELATIVE AMOUNTS OF NUCLEIC ACIDS IN A COMPLEX MIXTURE AND RETRIEVAL OF SPECIFIC SEQUENCES THEREFROM" to Kamb et al., issued May 9, 2000; US Patent No. 6,090,556 "METHOD FOR QUANTI-TATIVELY DETERMINING THE EXPRESSION OF A GENE" to Kato, issued July 18, 2000; and US Patent 6,040,138 "EXPRESSION MONITORING BY HYBRIDIZATION TO HIGH DENSITY OLIGONUCLEOTIDE ARRAYS" to Lockhart et al., issued March 21, 2000.

[0110] For example, cDNA inserts corresponding to candidates nucleotide sequences, in a standard TA cloning vector

are amplified by a polymerase chain reaction for approximately 30-40 cycles. The amplified PCR products are then arrayed onto a glass support by any of a variety of well known techniques, e.g., the VSLIPS™ technology described in US Patent No. 5,143,854. RNA, or cDNA corresponding to RNA, isolated from a subject sample of leukocytes is labeled, e.g., with a fluorescent tag, and a solution containing the RNA (or cDNA) is incubated under conditions favorable for hybridization, with the "probe" chip. Following incubation, and washing to eliminate non-specific hybridization, the labeled nucleic acid bound to the chip is detected qualitatively or quantitatively, and the resulting expression profile for the corresponding candidate nucleotide sequences is recorded. It is appreciated that the probe used for diagnostic purposes may be identical to the probe used during diagnostic nucleotide sequence discovery and validation. Alternatively, the probe sequence may be different than the sequence used in diagnostic nucleotide sequence discovery and validation. Multiple cDNAs from a nucleotide sequence that are non-overlapping or partially overlapping may also be used.

[0111] In another approach, oligonucleotides corresponding to members of an candidate nucleotide library are synthesized and spotted onto an array. Alternatively, oligonucleotides are synthesized onto the array using methods known in the art, e.g. Hughes, et al. *supra*. The oligonucleotide is designed to be complementary to any portion of the candidate nucleotide sequence. In addition, in the context of expression analysis for, e.g. diagnostic use of diagnostic nucleotide sets, an oligonucleotide can be designed to exhibit particular hybridization characteristics, or to exhibit a particular specificity and/or sensitivity, as further described below.

[0112] Hybridization signal may be amplified using methods known in the art, and as described herein, for example use of the Clontech kit (Glass Fluorescent Labeling Kit), Stratagene kit (Fairplay Microarray Labeling Kit), the Micromax kit (New England Nuclear, Inc.), the Genisphere kit (3DNA Submicro), linear amplification, e.g. as described in U.S. Patent No. 6,132,997 or described in Hughes, TR, et al., Nature Biotechnology, 19:343-347 (2001) and/or Westin et al. Nat Biotech. 18:199-204.

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[0113] Alternatively, fluorescently labeled cDNA are hybridized directly to the microarray using methods known in the art. For example, labeled cDNA are generated by reverse transcription using Cy3- and Cy5-conjugated deoxynucleotides, and the reaction products purified using standard methods. It is appreciated that the methods for signal amplification of expression data useful for identifying diagnostic nucleotide sets are also useful for amplification of expression data for diagnostic purposes.

[0114] Microarray expression may be detected by scanning the microarray with a variety of laser or CCD-based scanners, and extracting features with numerous software packages, for example, Imagene (Biodiscovery), Feature Extraction (Agilent), Scanalyze (Eisen, M. 1999. SCANALYZE User Manual; Stanford Univ., Stanford, CA. Ver 2.32.), GenePix (Axon Instruments).

[0115] In another approach, hybridization to microelectric arrays is performed, e.g. as described in Umek et al (2001) J Mol Diagn. 3:74-84. An affinity probe, e.g. DNA, is deposited on a metal surface. The metal surface underlying each probe is connected to a metal wire and electrical signal detection system. Unlabelled RNA or cDNA is hybridized to the array, or alternatively, RNA or cDNA sample is amplified before hybridization, e.g. by PCR. Specific hybridization of sample RNA or cDNA results in generation of an electrical signal, which is transmitted to a detector. See Westin (2000) Nat Biotech. 18:199-204 (describing anchored multiplex amplification of a microelectronic chip array); Edman (1997) NAR 25:4907-14; Vignali (2000) J Immunol Methods 243:243-55.

[0116] In another approach, a microfluidics chip is used for RNA sample preparation and analysis. This approach increases efficiency because sample preparation and analysis are streamlined. Briefly, microfluidics may be used to sort specific leukocyte sub-populations prior to RNA preparation and analysis. Microfluidics chips are also useful for, e.g., RNA preparation, and reactions involving RNA (reverse transcription, RT-PCR). Briefly, a small volume of whole, anti-coagulated blood is loaded onto a microfluidics chip, for example chips available from Caliper (Mountain View, CA) or Nanogen (San Diego, CA.) A microfluidics chip may contain channels and reservoirs in which cells are moved and reactions are performed. Mechanical, electrical, magnetic, gravitational, centrifugal or other forces are used to move the cells and to expose them to reagents. For example, cells of whole blood are moved into a chamber containing hypotonic saline, which results in selective lysis of red blood cells after a 20-minute incubation. Next, the remaining cells (leukocytes) are moved into a wash chamber and finally, moved into a chamber containing a lysis buffer such as guanidine isothyocyanate. The leukocyte cell lysate is further processed for RNA isolation in the chip, or is then removed for further processing, for example, RNA extraction by standard methods. Alternatively, the microfluidics chip is a circular disk containing ficoll or another density reagent. The blood sample is injected into the center of the disc, the disc is rotated at a speed that generates a centrifugal force appropriate for density gradient separation of mononuclear cells, and the separated mononuclear cells are then harvested for further analysis or processing.

[0117] It is understood that the methods of expression evaluation, above, although discussed in the context of discovery of diagnostic nucleotide sets, are also applicable for expression evaluation when using diagnostic nucleotide sets for, e.g. diagnosis of diseases, as further discussed below.

Evaluation of expression patterns

[0118] Expression patterns can be evaluated by qualitative and/or quantitative measures. Certain of the above described techniques for evaluating gene expression (as RNA or protein products) yield data that are predominantly qualitative in nature. That is, the methods detect differences in expression that classify expression into distinct modes without providing significant information regarding quantitative aspects of expression. For example, a technique can be described as a qualitative technique if it detects the presence or absence of expression of an candidate nucleotide sequence, i.e., an on/off pattern of expression. Alternatively, a qualitative technique measures the presence (and/or absence) of different alleles, or variants, of a gene product.

[0119] In contrast, some methods provide data that characterizes expression in a quantitative manner. That is, the methods relate expression on a numerical scale, e.g., a scale of 0-5, a scale of 1-10, a scale of + - +++, from grade 1 to grade 5, a grade from a to z, or the like. It will be understood that the numerical, and symbolic examples provided are arbitrary, and that any graduated scale (or any symbolic representation of a graduated scale) can be employed in the context of the present invention to describe quantitative differences in nucleotide sequence expression. Typically, such methods yield information corresponding to a relative increase or decrease in expression.

[0120] Any method that yields either quantitative or qualitative expression data is suitable for evaluating expression of candidate nucleotide sequence in a subject sample of leukocytes. In some cases, e.g., when multiple methods are employed to determine expression patterns for a plurality of candidate nucleotide sequences, the recovered data, e.g., the expression profile, for the nucleotide sequences is a combination of quantitative and qualitative data.

[0121] In some applications, expression of the plurality of candidate nucleotide sequences is evaluated sequentially. This is typically the case for methods that can be characterized as low- to moderate-throughput. In contrast, as the throughput of the elected assay increases, expression for the plurality of candidate nucleotide sequences in a sample or multiple samples of leukocytes, is assayed simultaneously. Again, the methods (and throughput) are largely determined by the individual practitioner, although, typically, it is preferable to employ methods that permit rapid, e.g. automated or partially automated, preparation and detection, on a scale that is time-efficient and cost-effective.

[0122] It is understood that the preceding discussion, while directed at the assessment of expression of the members of candidate libraries, is also applies to the assessment of the expression of members of diagnostic nucleotide sets, as further discussed below.

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[0123] In addition to, or in conjunction with the correlation of expression profiles and clinical data, it is often desirable to correlate expression patterns with the subject's genotype at one or more genetic loci. The selected loci can be, for example, chromosomal loci corresponding to one or more member of the candidate library, polymorphic alleles for marker loci, or alternative disease related loci (not contributing to the candidate library) known to be, or putatively associated with, a disease (or disease criterion). Indeed, it will be appreciated, that where a (polymorphic) allele at a locus is linked to a disease (or to a predisposition to a disease), the presence of the allele can itself be a disease criterion. [0124] Numerous well known methods exist for evaluating the genotype of an individual, including southern analysis, restriction fragment length polymorphism (RFLP) analysis, polymerase chain reaction (PCR), amplification length polymorphism (AFLP) analysis, single stranded conformation polymorphism (SSCP) analysis, single nucleotide polymorphism (SNP) analysis (e.g., via PCR, Taqman or molecular beacons), among many other useful methods. Many such procedures are readily adaptable to high throughput and/or automated (or semi-automated) sample preparation and analysis methods. Most, can be performed on nucleic acid samples recovered via simple procedures from the same sample of leukocytes as yielded the material for expression profiling. Exemplary techniques are described in, e.g., Sambrook, and Ausubel, supra.

Identification of the diagnostic nucleotide sets.

[0125] Identification of diagnostic nucleotide sets and disease specific target nucleotide sequence proceeds by correlating the leukocyte expression profiles with data regarding the subject's health status to produce a data set designated a "molecular signature." Examples of data regarding a patient's health status, also termed "disease criteria(ion)", is described below and in the Section titled "selected diseases," below. Methods useful for correlation analysis are further described elsewhere in the specification.

[0126] Generally, relevant data regarding the subject's health status includes retrospective or prospective health data, e.g., in the form of the subject's medical history, as provided by the subject, physician or third party, such as, medical diagnoses, laboratory test results, diagnostic test results, clinical events, or medication lists, as further described below. Such data may include information regarding a patient's response to treatment and/or a particular medication and data regarding the presence of previously characterized "risk factors." For example, cigarette smoking and obesity are pre-

viously identified risk factors for heart disease. Further examples of health status information, including diseases and disease criteria, is described in the section titled Selected diseases, below.

[0127] Typically, the data describes prior events and evaluations (i.e., retrospective data). However, it is envisioned that data collected subsequent to the sampling (i.e., prospective data) can also be correlated with the expression profile. The tissue sampled, e.g., peripheral blood, bronchial lavage, etc., can be obtained at one or more multiple time points and subject data is considered retrospective or prospective with respect to the time of sample procurement.

[0128] Data collected at multiple time points, called "longitudinal data", is often useful, and thus, the invention encompasses the analysis of patient data collected from the same patient at different time points. Analysis of paired samples, such as samples from a patient at different time, allows identification of differences that are specifically related to the disease state since the genetic variability specific to the patient is controlled for by the comparison. Additionally, other variables that exist between patients may be controlled for in this way, for example, the presence or absence of inflammatory diseases (e.g., rheumatoid arthritis) the use of medications that may effect leukocyte gene expression, the presence or absence of co-morbid conditions, etc. Methods for analysis of paired samples are further described below. Moreover, the analysis of a pattern of expression profiles (generated by collecting multiple expression profiles) provides information relating to changes in expression level over time, and may permit the determination of a rate of change, a trajectory, or an expression curve. Two longitudinal samples may provide information on the change in expression of a gene over time, while three longitudinal samples may be necessary to determine the "trajectory" of expression of a gene. Such information may be relevant to the diagnosis of a disease. For example, the expression of a gene may vary from individual, but a clinical event, for example, a heart attack, may cause the level of expression to double in each patient. In this example, clinically interesting information is gleaned from the change in expression level, as opposed to the absolute level of expression in each individual.

[0129] When a single patient sample is obtained, it may still be desirable to compare the expression profile of that sample to some reference expression profile. In this case, one can determine the change of expression between the patient's sample and a reference expression profile that is appropriate for that patient and the medical condition in question. For example, a reference expression profile can be determined for all patients without the disease criterion in question who have similar characteristics, such as age, sex, race, diagnoses etc.

[0130] Generally, small sample sizes of 20-100 samples are used to identify a diagnostic nucleotide set. Larger sample sizes are generally necessary to validate the diagnostic nucleotide set for use in large and varied patient populations, as further described below. For example, extension of gene expression correlations to varied ethnic groups, demographic groups, nations, peoples or races may require expression correlation experiments on the population of interest.

Expression Reference Standards

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[0131] Expression profiles derived from a patient (i.e., subjects diagnosed with, or exhibiting symptoms of, or exhibiting a disease criterion, or under a doctor's care for a disease) sample are compared to a control or standard expression RNA to facilitate comparison of expression profiles (e.g. of a set of candidate nucleotide sequences) from a group of patients relative to each other (i.e., from one patient in the group to other patients in the group, or to patients in another group).

[0132] The reference RNA used should have desirable features of low cost and simplicity of production on a large scale. Additionally, the reference RNA should contain measurable amounts of as many of the genes of the candidate library as possible.

[0133] For example, in one approach to identifying diagnostic nucleotide sets, expression profiles derived from patient samples are compared to a expression reference "Standard." Standard expression reference can be, for example, RNA derived from resting cultured leukocytes or commercially available reference RNA, such as Universal reference RNA from Stratagene. See Nature, V406, 8-17-00, p. 747-752. Use of an expression reference standard is particularly useful when the expression of large numbers of nucleotide sequences is assayed, e.g. in an array, and in certain other applications, e.g. qualitative PCR, RT-PCR, etc., where it is desirable to compare a sample profile to a standard profile, and/or when large numbers of expression profiles, e.g. a patient population, are to be compared. Generally, an expression reference standard should be available in largo quantities, should be a good substrate for amplification and labeling reactions, and should be capable of detecting a large percentage of candidate nucleic acids using suitable expression profiling technology.

[0134] Alternatively, or in addition, the expression profile derived from a patient sample is compared with the expression of an internal reference control gene, for example, β -actin or CD4. The relative expression of the profiled genes and the internal reference control gene (from the same individual) is obtained. An internal reference control may also be used with a reference RNA. For example, an expression profile for "gene 1" and the gene encoding CD4 can be determined in a patient sample and in a reference RNA. The expression of each gene can be expressed as the "relative" ratio of expression the gene in the patient sample compared with expression of the gene in the reference RNA. The expression ratio (sample/reference) for gene 1 may be divided by the expression ration for CD4 (sample/reference) and thus relative

expression of gene 1 to CD4 is obtained.

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[0135] The disclosure provides also a buffy coat control RNA useful for expression profiling, and a method of using control RNA produced from a population of buffy coat cells, the white blood cell layer derived from the centrifugation of whole blood. Buffy coat contains all white blood cells, including granulocytes, mononuclear cells and platelets. The disclosure provides also a method of preparing control RNA from buffy coat cells for use in expression profile analysis of leukocytes. Buffy coat fractions are obtained, e.g. from a blood bank or directly from individuals, preferably from a large number of individuals such that bias from individual samples is avoided and so that the RNA sample represents an average expression of a healthy population. Buffy coat fractions from about 50 or about 100, or more individuals are preferred. 10 ml buffy coat from each individual is used. Buffy coat samples are treated with an erthythrocyte lysis buffer, so that erthythrocytes are selectively removed. The leukocytes of the buffy coat layer are collected by centrifugation. Alternatively, the buffy cell sample can be further enriched for a particular leukocyte sub-populations, e.g. mononuclear cells, T-lymphocytes, etc. To enrich for mononuclear cells, the buffy cell pellet, above, is diluted in PBS (phosphate buffered saline) and loaded onto a non-polystyrene tube containing a polysucrose and sodium diatrizoate solution adjusted to a density of 1.077+/-0.001 g/ml. To enrich for T-lymphocytes, 45 ml of whole blood is treated with RosetteSep (Stem Cell Technologies), and incubated at room temperature for 20 minutes. The mixture is diluted with an equal volume of PBS plus 2% FBS and mixed by inversion. 30 ml of diluted mixture is layered on top of 15 ml DML medium (Stem Cell Technologies). The tube is centrifuged at 1200 x g, and the enriched cell layer at the plasma: medium interface is removed, washed with PBS + 2% FBS, and cells collected by centrifugation at 1200 x g. The cell pellet is treated with 5 ml of erythrocyte lysis buffer (EL buffer, Qiagen) for 10 minutes on ice, and enriched T-lymphoctes are collected by centrifugation.

[0136] In addition or alternatively, the buffy cells (whole buffy coat or sub-population, e.g. mononuclear fraction) can be cultured *in vitro* and subjected to stimulation with cytokines or activating chemicals such as phorbol esters or ionomycin. Such stimuli may increase expression of nucleotide sequences that are expressed in activated immune cells and might be of interest for leukocyte expression profiling experiments.

[0137] Following sub-population selection and/or further treatment, e.g. stimulation as described above, RNA is prepared using standard methods. For example, cells are pelleted and lysed with a phenol/guanidinium thiocyanate and RNA is prepared. RNA can also be isolated using a silica gel-based purification column or the column method can be used on RNA isolated by the phenol/guanidinium thiocyanate method. RNA from individual buffy coat samples can be pooled during this process, so that the resulting reference RNA represents the RNA of many individuals and individual bias is minimized or eliminated. In addition, a new batch of buffy coat reference RNA can be directly compared to the last batch to ensure similar expression pattern from one batch to another, using methods of collecting and comparing expression profiles described above/below. One or more expression reference controls are used in an experiment. For example, RNA derived from one or more of the following sources can be used as controls for an experiment: stimulated or unstimulated whole buffy coat, stimulated or unstimulated peripheral mononuclear cells, or stimulated or unstimulated T-lymphocytes.

[0138] Alternatively, the expression reference standard can be derived from any subject or class of subjects including healthy subjects or subjects diagnosed with the same or a different disease or disease criterion. Expression profiles from subjects in two distinct classes are compared to determine which subset of nucleotide sequences in the candidate library best distinguish between the two subject classes, as further discussed below. It will be appreciated that in the present context, the term "distinct classes" is relevant to at least one distinguishable criterion relevant to a disease of interest, a "disease criterion." The classes can, of course, demonstrate significant overlap (or identity) with respect to other disease criteria, or with respect to disease diagnoses, prognoses, or the like. The mode of discovery involves, e.g., comparing the molecular signature of different subject classes to each other (such as patient to control, patients with a first diagnosis to patients with a second diagnosis, etc.) or by comparing the molecular signatures of a single individual taken at different time points. The disclosure can be applied to a broad range of diseases, disease criteria, conditions and other clinical and/or epidemiological questions, as further discussed above/below.

[0139] It is appreciated that while the present discussion pertains to the use of expression reference controls while identifying diagnostic nucleotide sets, expression reference controls are also useful during use of diagnostic nucleotide sets, e.g. use of a diagnostic nucleotide set for diagnosis of a disease, as further described below.

Analysis of expression profiles

[0140] In order to facilitate ready access, e.g., for comparison, review, recovery, and/or modification, the molecular signatures/expression profiles are typically recorded in a database. Most typically, the database is a relational database accessible by a computational device, although other formats, e.g., manually accessible indexed files of expression profiles as photographs, analogue or digital imaging readouts, spreadsheets, etc. can be used. Further details regarding preferred embodiments are provided below. Regardless of whether the expression patterns initially recorded are analog or digital in nature and/or whether they represent quantitative or qualitative differences in expression, the expression

patterns, expression profiles (collective expression patterns), and molecular signatures (correlated expression patterns) are stored digitally and accessed via a database. Typically, the database is compiled and maintained at a central facility, with access being available locally and/or remotely.

[0141] As additional samples are obtained, and their expression profiles determined and correlated with relevant subject data, the ensuing molecular signatures are likewise recorded in the database. However, rather than each subsequent addition being added in an essentially passive manner in which the data from one sample has little relation to data from a second (prior or subsequent) sample, the algorithms optionally additionally query additional samples against the existing database to further refine the association between a molecular signature and disease criterion. Furthermore, the data set comprising the one (or more) molecular signatures is optionally queried against an expanding set of additional or other disease criteria. The use of the database in integrated systems and web embodiments is further described below.

Analysis of expression profile data from arrays

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[0142] Expression data is analyzed using methods well known in the art, including the software packages Imagene (Biodiscovery, Marina del Rey, CA), Feature Extraction Software (Agilent, Palo Alto, CA), and Scanalyze (Stanford University). In the discussion that follows, a "feature" refers to an individual spot of DNA on an array. Each gene may be represented by more than one feature. For example, hybridized microarrays are scanned and analyzed on an Axon Instruments scanner using GenePix 3.0 software (Axon Instruments, Union City, CA). The data extracted by GenePix is used for all downstream quality control and expression evaluation. The data is derived as follows. The data for all features flagged as "not found" by the software is removed from the dataset for individual hybridizations. The "not found" flag by GenePix indicates that the software was unable to discriminate the feature from the background. Each feature is examined to determine the value of its signal. The median pixel intensity of the background (Bn) is subtracted from the median pixel intensity of the feature (F_n) to produce the background-subtracted signal (hereinafter, "BGSS"). The BGSS is divided by the standard deviation of the background pixels to provide the signal-to-noise ratio (hereinafter, "S/N"). Features with a S/N of three or greater in both the Cy3 channel (corresponding to the sample RNA) and Cy5 channel (corresponding to the reference RNA) are used for further analysis (hereinafter denoted "useable features"). Alternatively, different S/Ns are used for selecting expression data for an analysis. For example, only expression data with signal to noise ratios > 3 might be used in an analysis. Alternatively, features with S/N values < 3 may be flagged as such and included in the analysis. Such flagged data sets include more values and may allow one to discover expression markers that would be missed otherwise. However, such data sets may have a higher variablilty than filtered data, which may decrease significance of findings or performance of correlation statistics.

[0143] For each usable feature (*i*), the expression level (*e*) is expressed as the logarithm of the ratio (R) of the Background Subtracted Signal (hereinafter "BGSS") for the Cy3 (sample RNA) channel divided by the BGSS for the Cy5 channel (reference RNA). This "log ratio" value is used for comparison to other experiments.

$$R_i = \frac{BGSS_{sample}}{BGSS_{reference}} \tag{0.1}$$

 $e_i = \log r_i \tag{0.2}$

[0144] Variation in signal across hybridizations may be caused by a number of factors affecting hybridization, DNA spotting, wash conditions, and labeling efficiency.

[0145] A single reference RNA may be used with all of the experimental RNAs, permitting multiple comparisons in addition to individual comparisons. By comparing sample RNAs to the same reference, the gene expression levels from each sample are compared across arrays, permitting the use of a consistent denominator for our experimental ratios.

[0146] Alternative methods of analyzing the data may involve 1) using the sample channel without normalization by the reference channel, 2) using an intensity-dependent normalization based on the reference which provides a greater correction when the signal in the reference channel is large, 3) using the data without background subtraction or subtracting an empirically derived function of the background intensity rather than the background itself.

Scaling

[0147] The data may be scaled (normalized) to control for labeling and hybridization variability within the experiment, using methods known in the art. Scaling is desirable because it facilitates the comparison of data between different

experiments, patients, etc. Generally the BGSS are scaled to a factor such as the median, the mean, the trimmed mean, and percentile. Additional methods of scaling include: to scale between 0 and 1, to subtract the mean, or to subtract the median.

[0148] Scaling is also performed by comparison to expression patterns obtained using a common reference RNA, as described in greater detail above. As with other scaling methods, the reference RNA facilitates multiple comparisons of the expression data, e.g., between patients, between samples, etc. Use of a reference RNA provides a consistent denominator for experimental ratios.

[0149] In addition to the use of a reference RNA, individual expression levels may be adjusted to correct for differences in labeling efficiency between different hybridization experiments, allowing direct comparison between experiments with different overall signal intensities, for example. A scaling factor (a) may be used to adjust individual expression levels as follows. The median of the scaling factor (a), for example, BGSS, is determined for the set of all features with a S/N greater than three. Next, the BGSS_i (the BGSS for each feature "i") is divided by the median for all features (a), generating a scaled ratio. The scaled ration is used to determine the expression value for the feature (e), or the log ratio.

$$S_i = \frac{BGSS_i}{a} \tag{0.3}$$

$$e_i = \log\left(\frac{Cy3S_i}{Cy5S_i}\right) \tag{0.4}$$

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[0150] In addition, or alternatively, control features are used to normalize the data for labeling and hybridization variability within the experiment. Control feature may be cDNA for genes from the plant, *Arabidopsis thaliana*, that are included when spotting the mini-array. Equal amounts of RNA complementary to control cDNAs are added to each of the samples before they were labeled. Using the signal from these control genes, a normalization constant (*L*) is determined according to the following formula:

$$L_{j} = \frac{\sum_{i=1}^{N} BGSS_{j,i}}{N}$$

$$\sum_{j=1}^{K} \sum_{i=1}^{N} BGSS_{j,i}}$$

$$K$$

where BGSS_i is the signal for a specific feature, N is the number of A. thaliana control features, K is the number of hybridizations, and L_i is the normalization constant for each individual hybridization.

[0151] Using the formula above, the mean for all control features of a particular hybridization and dye (e.g., Cy3) is calculated. The control feature means for all Cy3 hybridizations are averaged, and the control feature mean in one hybridization divided by the average of all hybridizations to generate a normalization constant for that particular Cy3 hybridization (L_j), which is used as a in equation (0.3). The same normalization steps may be performed for Cy3 and Cy5 values.

[0152] An alternative scaling method can also be used. The log of the ratio of Green/Red is determined for all features. The median log ratio value for all features is determined. The feature values are then scaled using the following formula: Log_Scaled_Feature_Ratio = Log_Feature_Ratio - Median_Log_Ratio.

[0153] Many additional methods for normalization exist and can be applied to the data. In one method, the average ratio of Cy3 BGSS / Cy5 BGSS is determined for all features on an array. This ratio is then scaled to some arbitrary number, such as 1 or some other number. The ratio for each probe is then multiplied by the scaling factor required to bring the average ratio to the chosen level. This is performed for each array in an analysis. Alternatively, the ratios are normalized to the average ratio across all arrays in an analysis. Other methods of normalization include forcing the distribution of signal strengths of the various arrays into greater agreement by transforming them to match certain points (quartiles, or deciles, etc.) in a standard distribution, or in the most extreme case using the rank of the signal of each oligonucleotide relative to the other oligonucleotides on the array.

[0154] If multiple features are used per gene sequence or oligonucleotide, these repeats can be used to derive an average expression value for each gene. If some of the replicate features are of poor qualitay and don't meet requirements for analysis, the remaining features can be used to represent the gene or gene sequence.

Correlation analysis

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[0155] Correlation analysis is performed to determine which array probes have expression behavior that best distinguishes or serves as markers for relevant groups of samples representing a particular clinical condition. Correlation analysis, or comparison among samples representing different disease criteria (e.g., clinical conditions), is performed using standard statistical methods. Numerous algorithms are useful for correlation analysis of expression data, and the selection of algorithms depends in part on the data analysis to be performed. For example, algorithms can be used to identify the single most informative gene with expression behavior that reliably classifies samples, or to identify all the genes useful to classify samples. Alternatively, algorithms can be applied that determine which set of 2 or more genes have collective expression behavior that accurately classifies samples. The use of multiple expression markers for diagnostics may overcome the variability in expression of a gene between individuals, or overcome the variability intrinsic to the assay. Multiple expression markers may include redundant markers (surrogates), in that two or more genes or probes may provide the same information with respect to diagnosis. This may occur, for example, when two or more genes or gene probes are coordinately expressed. For diagnostic application, it may be appropriate to utilize a gene and one or more of its surrogates in the assay. This redundancy may overcome failures (technical or biological) of a single marker to distinguish samples. Alternatively, one or more surrogates may have properties that make them more suitable for assay development, such as a higher baseline level of expression, better cell specificity, a higher fold change between sample groups or more specific sequence for the design of PCR primers or complimentary probes. It will be appreciated that while the discussion above pertains to the analysis of RNA expression profiles the discussion is equally applicable to the analysis of profiles of proteins or other molecular markers.

[0156] Prior to analysis, expression profile data may be formatted or prepared for analysis using methods known in the art. For example, often the log ratio of scaled expression data for every array probe is calculated using the following formula:

log (Cy 3 BGSS/ Cy5 BGSS),

where Cy 3 signal corresponds to the expression of the gene in the clinical sample, and Cy5 signal corresponds to expression of the gene in the reference RNA.

[0157] Data may be further filtered depending on the specific analysis to be done as noted below. For example, filtering may be aimed at selecting only samples with expression above a certain level, or probes with variability above a certain level between sample sets.

[0158] The following non-limiting discussion consider several statistical methods known in the art. Briefly, the t-test and ANOVA are used to identify single genes with expression differences between or among populations, respectively. Multivariate methods are used to identify a set of two or more genes for which expression discriminates between two disease states more specifically than expression of any single gene.

t-test

[0159] The simplest measure of a difference between two groups is the Student's t test. See, e.g., Welsh et al. (2001) Proc Natl Acad Sci USA 98:1176-81 (demonstrating the use of an unpaired Student's t-test for the discovery of differential gene expression in ovarian cancer samples and control tissue samples). The t- test assumes equal variance and normally distributed data. This test identifies the probability that there is a difference in expression of a single gene between two groups of samples. The number of samples within each group that is required to achieve statistical significance is dependent upon the variation among the samples within each group. The standard formula for a t-test is:

$t(e_i) = \frac{\overline{e}_{i,c} - \overline{e}_{i,t}}{\sqrt{(s_{i,c}^2/n_c) + (s_{i,t}^2/n_t)}},$ (0.5)

where $\overline{e_i}$ is the difference between the mean expression level of gene i in groups c and t, $s_{i,c}$ is the variance of gene x in group c and $s_{i,t}$ is the variance of gene x in group t. n_c and n_t are the numbers of samples in groups c and t.

[0160] The combination of the t statistic and the degrees of freedom [min(n_t , n_c)-1] provides a p value, the probability of rejecting the null hypothesis. A p-value of \leq 0.01, signifying a 99 percent probability the mean expression levels are different between the two groups (a 1% chance that the mean expression levels are in fact not different and that the observed difference occurred by statistical chance), is often considered acceptable.

[0161] When performing tests on a large scale, for example, on a large dataset of about 8000 genes, a correction factor must be included to adjust for the number of individual tests being performed. The most common and simplest correction is the Bonferroni correction for multiple tests, which divides the p-value by the number of tests run. Using this test on an 8000 member dataset indicates that a p value of ≤ 0.00000125 is required to identify genes that are likely to be truly different between the two test conditions.

Significance analysis for microarrays (SAM)

[0162] Significance analysis for microarrays (SAM) (Tusher 2001) is a method through which genes with a correlation between their expression values and the response vector are statistically discovered and assigned a statistical significance. The ratio of false significant to significant genes is the False Discovery Rate (FDR). This means that for each threshold there are a set of genes which are called significant, and the FDR gives a confidence level for this claim. If a gene is called differentially expressed between 2 classes by SAM, with a FDR of 5%, there is a 95% chance that the gene is actually differentially expressed between the classes. SAM takes intoaccount the variability and large number of variables of microarrays. SAM will identiy genes that are most globally differentially expressed between the classes. Thus, important genes for identifying and classifying outlier samples or patients may not be identified by SAM.

Non-Parametric Tests

[0163] Wilcoxon's signed ranks method is one example of a non-parametric test and is utilized for paired comparisons. See e.g., Sokal and Rohlf (1987) Introduction to Biostatistics 2nd edition, WH Freeman, New York. At least 6 pairs are necessary to apply this statistic. This test is useful for analysis of paired expression data (for example, a set of patients who have cardiac transplant biopsy on 2 occasions and have a grade 0 on one occasion and a grade 3A on another). The Fisher Exact Test with a threshold and the Mann-Whitney Test are other non-parametric tests that may be used.

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[0164] Differences in gene expression across multiple related groups may be assessed using an Analysis of Variance (ANOVA), a method well known in the art (Michelson and Schofield, 1996).

35 Multivariate analysis

[0165] Many algorithms suitable for multivariate analysis are known in the art. Generally, a set of two or more genes for which expression discriminates between two disease states more specifically than expression of any single gene is identified by searching through the possible combinations of genes using a criterion for discrimination, for example the expression of gene X must increase from normal 300 percent, while the expression of genes Y and Z must decrease from normal by 75 percent. Ordinarily, the search starts with a single gene, then adds the next best fit at each step of the search. Alternatively, the search starts with all of the genes and genes that do not aid in the discrimination are eliminated step-wise.

45 Paired samples

[0166] Paired samples, or samples collected at different time-points from the same patient, are often useful, as described above. For example, use of paired samples permits the reduction of variation due to genetic variation among individuals. In addition, the use of paired samples has a statistical significance, in that data derived from paired samples can be calculated in a different manner that recognizes the reduced variability. For example, the formula for a t-test for paired samples is:

$$t(e_x) = \frac{\overline{D}_{\bar{e}_x}}{\sqrt{\frac{\sum D^2 - (\sum D)^2 / b}{b - 1}}},$$
(0.5)

where D is the difference between each set of paired samples and b is the number of sample pairs. \overline{D} is the mean of the differences between the members of the pairs. In this test, only the differences between the paired samples are considered, then grouped together (as opposed to taking all possible differences between groups, as would be the case with an ordinary t-test). Additional statistical tests useful with paired data, e.g., ANOVA and Wilcoxon's signed rank test, are discussed above.

Diagnostic classification

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[0167] Once a discriminating set of genes is identified, the diagnostic classifier (a mathematical function that assigns samples to diagnostic categories based on expression data) is applied to unknown sample expression levels.

[0168] Methods that can be used for this analysis include the following non-limiting list:

CLEAVER is an algorithm used for classification of useful expression profile data. See Raychaudhuri et al. (2001) Trends Biotechnol 19:189-193. CLEAVER uses positive training samples (e.g., expression profiles from samples known to be derived from a particular patient or sample diagnostic category, disease or disease criteria), negative training samples (e.g., expression profiles from samples known not to be derived from a particular patient or sample diagnostic category, disease or disease criteria) and test samples (e.g., expression profiles obtained from a patient), and determines whether the test sample correlates with the particular disease or disease criteria, or does not correlate with a particular disease or disease criteria. CLEAVER also generates a list of the 20 most predictive genes for classification.

[0169] Artificial neural networks (hereinafter, "ANN") can be used to recognize patterns in complex data sets and can discover expression criteria that classify samples into more than 2 groups. The use of artificial neural networks for discovery of gene expression diagnostics for cancers using expression data generated by oligonucleotide expression microarrays is demonstrated by Khan et al. (2001) Nature Med. 7:673-9. Khan found that 96 genes provided 0% error rate in classification of the tumors. The most important of these genes for classification was then determined by measuring the sensitivity of the classification to a change in expression of each gene. Hierarchical clustering using the 96 genes results in correct grouping of the cancers into diagnostic categories.

[0170] Golub uses cDNA microarrays and a distinction calculation to identify genes with expression behavior that distinguishes myeloid and lymphoid leukemias. See Golub et al. (1999) Science 286:531-7. Self organizing maps were used for new class discovery. Cross validation was done with a "leave one out" analysis. 50 genes were identified as useful markers. This was reduced to as few as 10 genes with equivalent diagnostic accuracy.

[0171] Hierarchical and non-hierarchical clustering methods are also useful for identifying groups of genes that correlate with a subset of clinical samples such as with transplant rejection grade. Alizadeh used hierarchical clustering as the primary tool to distinguish different types of diffuse B-cell lymphomas based on gene expression profile data. See Alizadeh et al. (2000) Nature 403:503-11. Alizadeh used hierarchical clustering as the primary tool to distinguish different types of diffuse B-cell lymphomas based on gene expression profile data. A cDNA array carrying 17856 probes was used for these experiments, 96 samples were assessed on 128 arrays, and a set of 380 genes was identified as being useful for sample classification.

[0172] Perou demonstrates the use of hierarchical clustering for the molecular classification of breast tumor samples based on expression profile data. See Perou et al. (2000) Nature 406:747-52. In this work, a cDNA array carrying 8102 gene probes was used. 1753 of these genes were found to have high variation between breast tumors and were used for the analysis.

[0173] Hastie describes the use of gene shaving for discovery of expression markers. Hastie et al. (2000) Genome Biol. 1(2):RESEARCH 0003.1-0003.21. The gene shaving algorithm identifies sets of genes with similar or coherent expression patterns, but large variation across conditions (RNA samples, sample classes, patient classes). In this manner, genes with a tight expression pattern within a transplant rejection grade, but also with high variability across rejection grades are grouped together. The algorithm takes advantage of both characteristics in one grouping step. For example, gene shaving can identify useful marker genes with co-regulated expression. Sets of useful marker genes can be reduced to a smaller set, with each gene providing some non-redundant value in classification. This algorithm was used on the data set described in Alizadeh et al., supra, and the set of 380 informative gene markers was reduced to 234.

[0174] Supervised harvesting of expression trees (Hastie 2001) identifies genes or clusters that best distinguish one class from all the others on the data set. The method is used to identify the genes/clusters that can best separate one class versus all the others for datasets that include two or more classes or all classes from each other. This algorithm can be used for discovery or testing of a diagnostic gene set.

[0175] CART is a decision tree classification algorithm (Breiman 1984). From gene expression and or other data, CART can develop a decision tree for the classification of samples. Each node on the decision tree involves a query about the expression level of one or more genes or variables. Samples that are above the threshold go down one branch

of the decision tree and samples that are not go down the other branch. See example 4 for further description of its use in classification analysis and examples of its usefulness in discovering and implementing a diagnostic gene set. CART identifies surrogates for each splitter (genes that are the next best substitute for a useful gene in classification.

[0176] Multiple Additive Regression Trees (Friedman, JH 1999, MART) is similar to CART in that it is a classification algorithm that builds decision trees to distinguish groups. MART builds numerous trees for any classification problem and the resulting model involves a combination of the multiple trees. MART can select variables as it build models and thus can be used on large data sets, such as those derived from an 8000 gene microarray. Because MART uses a combination of many trees and does not take too much information from any one tree, it resists over training. MART identifies a set of genes and an algorithm for their use as a classifier.

[0177] A Nearest Shrunken Centroids Classifier can be applied to microarray or other data sets by the methods described by Tibshirani et al. 2002. This algorithms also identified gene sets for classification and determines their 10 fold cross validation error rates for each class of samples. The algorithm determines the error rates for models of any size, from one gene to all genes in the set. The error rates for either or both sample classes can are minimized when a particular number of genes are used. When this gene number is determined, the algorithm associated with the selected genes can be identified and employed as a classifier on prospective sample.

[0178] Once a set of genes and expression criteria for those genes have been established for classification, cross validation is done. There are many approaches, including a 10 fold cross validation analysis in which 10% of the training samples are left out of the analysis and the classification algorithm is built with the remaining 90%. The 10% are then used as a test set for the algorithm. The process is repeated 10 times with 10% of the samples being left out as a test set each time. Through this analysis, one can derive a cross validation error which helps estimate the robustness of the algorithm for use on prospective (test) samples.

[0179] Clinical data are gathered for every patient sample used for expression analysis. Clinical variables can be quantitative or non-quantitative. A clinical variable that is quantitative can be used as a variable for significance or classification analysis. Non-quantitative clinical variables, such as the sex of the patient, can also be used in a significance analysis or classification analysis with some statistical tool. It is appreciated that the most useful diagnostic gene set for a condition may be optimal when considered along with one or more predictive clinical variables. Clinical data can also be used as supervising vectors for a correlation analysis. That is to say that the clinical data associated with each sample can be used to divide the samples into meaningful diagnostic categories for analysis. For example, samples can be divided into 2 or more groups based on the presence or absence of some diagnostic criterion (a). In addition, clinical data can be utilized to select patients for a correlation analysis or to exclude them based on some undesirable characteristic, such as an ongoing infection, a medicine or some other issue. Clinical data can also be used to assess the pretest probability of an outcome. For example, patients who are female are much more likely to be diagnosed as having systemic lupus erythematosis than patients who are male.

[0180] Once a set of genes are identified that classify samples with acceptable accuracy. These genes are validated as a set using new samples that were not used to discover the gene set. These samples can be taken from frozen archieves from the discovery clinical study or can be taken from new patients prospectively. Validation using a "test set" of samples can be done using expression profiling of the gene set with microarrays or using real-time PCR for each gene on the test set samples. Alternatively, a different expression profiling technology can be used.

Immune Monitoring

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[0181] Leukocyte gene expression can be used to monitor the immune system. Immune monitoring examines both the level of gene expression for a set of genes in a given cell type and for genes which are expressed in a cell type selective manner gene expression monitoring will also detect the presence or absence of new cell types, progenitor cells, differentiation of cells and the like. Gene expression patterns may be associated with activation or the resting state of cells of the immune system that are responsible for or responsive to a disease state. For example, in the process of transplant rejection, cells of the immune system are activated by the presence of the foreign tissue. Genes and gene sets that monitor and diagnose this process are providing a measure of the level and type of activation of the immune system. Genes and gene sets that are useful in monitoring the immune system may be useful for diagnosis and monitoring of all diseases that involve the immune system. Some examples are transplant rejection, rheumatoid arthritis, lupus, inflammatory bowel diseases, multiple sclerosis, HIV/AIDS, and viral, bacterial and fungal infection. All disorders and diseases disclosed herein are contemplated. Genes and gene sets that monitor immune activation are useful for monitoring response to immunosuppressive drug therapy, which is used to decrease immune activation. Genes are found to correlate with immune activation by correlation of expression patterns to the known presence of immune activation or quiescence in a sample as determined by some other test.

Selected Diseases

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[0182] In principle, diagnostic nucleotide sets of the invention may be developed and applied to essentially any disease, or disease criterion, as long as at least one subset of nucleotide sequences is differentially expressed in samples derived from one or more individuals with a disease criteria or disease and one or more individuals without the disease criteria or disease, wherein the individual may be the same individual sampled at different points in time, or the individuals may be different individuals (or populations of individuals). For example, the subset of nucleotide sequences may be differentially expressed in the sampled tissues of subjects with the disease or disease criterion (e.g., a patient with a disease or disease criteria) as compared to subjects without the disease or disease criterion (e.g., patients without a disease (control patients)). Alternatively, or in addition, the subset of nucleotide sequence(s) may be differentially expressed in different samples taken from the same patient, e.g at different points in time, at different disease stages, before and after a treatment, in the presence or absence of a risk factor, etc.

[0183] Expression profiles corresponding to sets of nucleotide sequences that correlate not with a diagnosis, but rather with a particular aspect of a disease can also be used to identify the diagnostic nucleotide sets and disease specific target nucleotide sequences of the invention. For example, such an aspect, or disease criterion, can relate to a subject's medical or family history, e.g., childhood illness, cause of death of a parent or other relative, prior surgery or other intervention, medications, symptoms (including onset and/or duration of symptoms), etc. Alternatively, the disease criterion can relate to a diagnosis, e.g., hypertension, diabetes, atherosclerosis, or prognosis (e.g., prediction of future diagnoses, events or complications), e.g., acute myocardial infarction, restenosis following angioplasty, reperfusion injury, allograft rejection, rheumatoid arthritis or systemic lupus erythematosis disease activity or the like. In other cases, the disease criterion corresponds to a therapeutic outcome, e.g., transplant rejection, bypass surgery or response to a medication, restenosis after stent implantation, collateral vessel growth due to therapeutic angiogenesis therapy, decreased angina due to revascularization, resolution of symptoms associated with a myriad of therapies, and the like. Alternatively, the disease criteria corresponds with previously identified or classic risk factors and may correspond to prognosis or future disease diagnosis. As indicated above, a disease criterion can also correspond to genotype for one or more loci. Disease criteria (including patient data) may be collected (and compared) from the same patient at different points in time, from different patients, between patients with a disease (criterion) and patients respresenting a control population, etc. Longitudinal data, i.e., data collected at different time points from an individual (or group of individuals) may be used for comparisons of samples obtained from an individual (group of individuals) at different points in time, to permit identification of differences specifically related to the disease state, and to obtain information relating to the change in expression over time, including a rate of change or trajectory of expression over time. The usefulness of longitudinal data is further discussed in the section titled "Identification of diagnostic nucleotide sets of the invention".

[0184] It is further understood that diagnostic nucleotide sets may be developed for use in diagnosing conditions for which there is no present means of diagnosis. For example, in rheumatoid arthritis, joint destruction is often well under way before a patient experience symptoms of the condition. A diagnostic nucleotide set may be developed that diagnoses rheumatic joint destruction at an earlier stage than would be possible using present means of diagnosis, which rely in part on the presentation of symptoms by a patient. Diagnostic nucleotide sets may also be developed to replace or augment current diagnostic procedures. For example, the use of a diagnostic nucleotide set to diagnose cardiac allograft rejection may replace the current diagnostic test, a graft biopsy.

[0185] It is understood that the following discussion of diseases is exemplary and non-limiting, and further that the general criteria discussed above, e.g. use of family medical history, are generally applicable to the specific diseases discussed below.

[0186] In addition to leukocytes, as described throughout, the general method is applicable to nucleotide sequences that are differentially expressed in any subject tissue or cell type, by the collection and assessment of samples of that tissue or cell type. However, in many cases, collection of such samples presents significant technical or medical problems given the current state of the art.

Organ transplant rejection and success

[0187] A frequent complication of organ transplantation is recognition of the transplanted organ as foreign by the immune system resulting in rejection. Diagnostic nucleotide sets can be identified and validated for monitoring organ transplant success, rejection and treatment. Medications currently exist that suppress the immune system, and thereby decrease the rate of and severity of rejection. However, these drugs also suppress the physiologic immune responses, leaving the patient susceptible to a wide variety of opportunistic infections and cancers. At present there is no easy, reliable way to diagnose transplant rejection. Organ biopsy is the preferred method, but this is expensive, painful and associated with significant risk and has inadequate sensitivity for focal rejection.

[0188] Diagnostic nucleotide sets of the present disclosure can be developed and validated for use as diagnostic tests for transplant rejection and success. It is appreciated that the methods of identifying diagnostic nucleotide sets are

applicable to any organ transplant population. For example, diagnostic nucleotide sets are developed for cardiac allograft rejection and success.

[0189] In some cases, disease criteria correspond to acute stage rejection diagnosis based on organ biopsy and graded using the International Society for Heart and Lung Transplantation ("ISHLT") criteria. This grading system classifies endomyocardial biopsies on the histological level as Grade 0, 1A, 1B, 2, 3A, 3B, or 4. Grade 0 biopsies have no evidence of rejection, while each successive grade has increased severity of leukocyte infiltration and/or damage to the graft myocardial cells. It is appreciated that there is variability in the Grading systems between medical centers and pathologists and between repeated readings of the same pathologist at different times. When using the biopsy grade as a disease criterion for leukocyte gene expression correlation analysis, it may be desirable to have a single pathologist read all biopsy slides or have multiple pathologists read all slides to determine the variability in this disease criterion. It is also appreciated that cardiac biopsy, in part due to variability, is not 100% sensitive or 100% specific for diagnosing acute rejection. When using the cardiac biopsy grade as a disease criterion for the discovery of diagnostic gene sets, it may be desirable to divide patient samples into diagnostic categories based on the grades. Examples of such classes are those patients with: Grade 0 vs. Grades 1A-4, Grade 0 vs. Grades 1B-4, Grade 0 vs. Grades 2-4, Grade 0-1 vs. Grade 3A-4, or Grade 0 vs. Grade 3A-4.

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[0190] Other disease criteria correspond to the cardiac biopsy results <u>and</u> other criteria, such as the results of cardiac function testing by echocardiography, hemodynamics assessment by cardiac catheterization, CMV infection, weeks post transplant, medication regimen, demographics and/or results of other diagnostic tests.

[0191] Other disease criteria correspond to information from the patient's medical history and information regarding the organ donor. Alternatively, disease criteria include the presence or absence of cytomegalovirus (CMV) infection, Epstein-Barr virus (EBV) infection, allograft dysfunction measured by physiological tests of cardiac function (e.g., hemodynamic measurements from catheterization or echocardiograph data), and symptoms of other infections. Alternatively, disease criteria correspond to therapeutic outcome, e.g. graft failure, re-transplantation, death, hospitalization, need for intravenous immunosuppression, transplant vasculopathy, response to immunosuppressive medications, etc. Disease criteria may further correspond to a rejection episode of at least moderate histologic grade, which results in treatment of the patient with additional corticosteroids, anti-T cell antibodies, or total lymphoid irradiation; a rejection with histologic grade 2 or higher; a rejection with histologic grade <2; the absence of histologic rejection and normal or unchanged allograft function (based on hemodynamic measurements from catheterization or on echocardiographic data); the presence of severe allograft dysfunction or worsening allograft dysfunction during the study period (based on hemodynamic measurements from catheterization or on echocardiographic data).; documented CMV infection by culture, histology, or PCR, and at least one clinical sign or symptom of infection; specific graft biopsy rejection grades; rejection of mild to moderate histologic severity prompting augmentation of the patient's chronic immunosuppressive regimen; rejection of mild to moderate severity with allograft dysfunction prompting plasmaphoresis or a diagnosis of "humoral" rejection; infections other than CMV, especially infection with Epstein Barr virus (EBV); lymphoproliferative disorder (also called post-transplant lymphoma); transplant vasculopathy diagnosed by increased intimal thickness on intravascular ultrasound (IVUS), angiography, or acute myocardial infarction; graft failure or retransplantation; and all cause mortality. Further specific examples of clinical data useful as disease criteria are provided in Example 3.

[0192] In another example, diagnostic nucleotide sets are developed and validated for use in diagnosis and monitoring of kidney allograft recipients. Disease criteria correspond to, e.g., results of biopsy analysis for kidney allograft rejection, serum creatine level, creatinine clearance, radiological imaging results for the kidney and urinalysis results. Another disease criterion corresponds to the need for hemodialysis, retransplantation, death or other renal replacement therapy. Diagnostic nucleotide sets are developed and validated for use in diagnosis and treatment of bone marrow transplant and liver transplantation pateints, respectively. Disease criteria for bone marrow transplant correspond to the diagnosis and monitoring of graft rejection and/or graft versus host disease, the recurrence of cancer, complications due to immunosuppression, hematologic abnormalities, infection, hospitalization and/or death. Disease criteria for liver transplant rejection include levels of serum markers for liver damage and liver function such as AST (aspartate aminotransferase), ALT (alanine aminotransferase), Alkaline phosphatase, GGT, (gamma-glutamyl transpeptidase) Bilirubin, Albumin and Prothrombin time. Further disease criteria correspond to hepatic encephalopathy, medication usage, ascites, graft failure, retransplantation, hospitalization, complications of immunosuppression, results of diagnostic tests, results of radiological testing, death and histological rejection on graft biopsy. In addition, urine can be utilized for at the target tissue for profiling in renal transplant, while biliary and intestinal secretions and feces may be used favorably for hepatic or intestinal organ allograft rejection. Diagnostic nuclotide sets can also be discovered and developed for the diagnosis and monitoring of chronic renal allograft rejection.

[0193] In the case of renal allografts, gene expression markers may be identified that are secreted proteins. These proteins may be detected in the urine of allograft recipients using standard immunoassays. Proteins are more likely to be present in the urine if they are of low molecular weight. Lower molecular weight proteins are more likely to pass through the glomerular membrane and into the urine.

[0194] In another example, diagnostic nucleotide sets are developed and validated for use in diagnosis and treatment

of xenograft recipients. This can include the transplantation of any organ from a non-human animal to a human or between non-human animals. Considerations for discovery and application of diagnostics and therapeutics and for disease criterion are substantially similar to those for allograft transplantation between humans.

[0195] In another example, diagnostic nucleotide sets are developed and validated for use in diagnosis and treatment of artificial organ recipients. This includes, but is not limited to mechanical circulatory support, artificial hearts, left ventricular assist devices, renal replacement therapies, organ prostheses and the like. Disease criteria are thrombosis (blood clots), infection, death, hospitalization, and worsening measures of organ function (e.g., hemodynamics, creatinine, liver function testing, renal function testing, functional capacity).

[0196] In another example, diagnostic nucleotide sets are developed and validated for use in matching donor organs to appropriate recipients. Diagnostic gene set can be discovered that correlate with successful matching of donor organ to recipient. Disease criteria include graft failure, acute and chronic rejection, death, hospitalization, immunosuppressive drug use, and complications of immunosuppression. Gene sets may be assayed from the donor or recipient's peripheral blood, organ tissue or some other tissue.

[0197] In another example, diagnostic nucleotide sets are developed and validated for use in diagnosis and induction of patient immune tolerance (decrease rejection of an allograft by the host immune system). Disease criteria include rejection, assays of immune activation, need for immunosupression and all disease criteria noted above for transplantation of each organ.

Viral diseases

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[0198] Diagnostic leukocyte nucleotide sets may be developed and validated for use in diagnosing viral disease, as well as diagnosing and monitoring transplant rejection. In another aspect, viral nucleotide sequences may be added to a leukocyte nucleotide set for use in diagnosis of viral diseases, as well as diagnosing and monitoring transplant rejection. Alternatively, viral nucleotide sets and leukocyte nucleotides sets may be used sequentially.

Epstein-Barr virus (EBV)

[0199] EBV causes a variety of diseases such as mononucleosis, B-cell lymphoma, and pharyngeal carcinoma. It infects mononuclear cells and circulating atypical lymphocytes are a common manifestation of infection. Peripheral leukocyte gene expression is altered by infection. Transplant recipients and patients who are immunosuppressed are at increased risk for EBV-associated lymphoma.

[0200] Diagnostic nucleotide sets may be developed and validated for use in diagnosis and monitoring of EBV, as well as diagnosing and monitoring transplant rejection. In one aspect, the diagnostic nucleotide set is a leukocyte nucleotide set. Alternatively, EBV nucleotide sequences are added to a leukocyte nucleotide set, for use in diagnosing EBV. Disease criteria correspond with diagnosis of EBV, and, in patients who are EBV-sero-positive, presence (or prospective occurrence) of EBV-related illnesses such as mononucleosis, and EBV-associated lymphoma. Diagnostic nucleotide sets are useful for diagnosis of EBV, and prediction of occurrence of EBV-related illnesses.

Cytomegalovirus (CMV)

[0201] Cytomegalovirus cause inflammation and disease in almost any tissue, particularly the colon, lung, bone marrow and retina, and is a very important cause of disease in immunosuppressed patients, e.g. transplant, cancer, AIDS. Many patients are infected with or have been exposed to CMV, but not all patients develop clinical disease from the virus. Also, CMV negative recipients of allografts that come from CMV positive donors are at high risk for CMV infection. As immunosuppressive drugs are developed and used, it is increasingly important to identify patients with current or impending clinical CMV disease, because the potential benefit of immunosuppressive therapy must be balanced with the increased rate of clinical CMV infection and disease that may result from the use of immunosuppression therapy. CMV may also play a role in the occurrence of atherosclerosis or restenosis after angioplasty. CMV expression also correlates to transplant rejection, and is useful in diagnosing and monitoring transplant rejection.

[0202] Diagnostic nucleotide sets are developed for use in diagnosis and monitoring of CMV infection or re-activation of CMV infection. In one aspect, the diagnostic nucleotide set is a leukocyte nucleotide set. In another aspect, CMV nucleotide sequences are added to a leukocyte nucleotide set, for use in diagnosing CMV. Disease criteria correspond to diagnosis of CMV (e.g., sero-positive state) and presence of clinically active CMV. Disease criteria may also correspond to prospective data, e.g. the likelihood that CMV will become clinically active or impending clinical CMV infection. Antiviral medications are available and diagnostic nucleotide sets can be used to select patients for early treatment, chronic suppression or prophylaxis of CMV activity.

Hepatitis Band C

[0203] These chronic viral infections affect about 1.25 and 2.7 million patients in the US, respectively. Many patients are infected, but suffer no clinical manifestations. Some patients with infection go on to suffer from chronic liver failure, cirrhosis and hepatic carcinoma.

[0204] Diagnostic nucleotide sets are developed for use in diagnosis and monitoring of HBV or HCV infection. In one aspect, the diagnostic nucleotide set is a leukocyte nucleotide set. In another aspect, viral nucleotide sequences are added to a leukocyte nucleotide set, for use in diagnosing the virus and monitoring progression of liver disease. Disease criteria correspond to diagnosis of the virus (e.g., sero-positive state or other disease symptoms). Alternatively, disease criteria correspond to liver damage, e.g., elevated alkaline phosphatase, ALT, AST or evidence of ongoing hepatic damage on liver biopsy. Alternatively, disease criteria correspond to serum liver tests (AST, ALT, Alkaline Phosphatase, GGT, PT, bilirubin), liver biopsy, liver ultrasound, viral load by serum PCR, cirrhosis, hepatic cancer, need for hospitalization or listing for liver transplant. Diagnostic nucleotide sets are used to diagnose HBV and HCV, and to predict likelihood of disease progression. Antiviral therapeutic usage, such as Interferon gamma and Ribavirin, can also be disease criteria.

HIV

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[0205] HIV infects T cells and certainly causes alterations in leukocyte expression. Diagnostic nucleotide sets are developed for diagnosis and monitoring of HIV. In one aspect, the diagnostic nucleotide set is a leukocyte nucleotide set. In another aspect, viral nucleotide sequences are added to a leukocyte nucleotide set, for use in diagnosing the virus. Disease criteria correspond to diagnosis of the virus (e.g., sero-positive state). In addition, disease criteria correspond to viral load, CD4 T cell counts, opportunistic infection, response to antiretroviral therapy, progression to AIDS, rate of progression and the occurrence of other HIV related outcomes (e.g., malignancy, CNS disturbance). Response to antiretrovirals may also be disease criteria.

Pharmacogenomics

[0206] Pharmocogenomics is the study of the individual propensity to respond to a particular drug therapy (combination of therapies). In this context, response can mean whether a particular drug will work on a particular patient, e.g. some patients respond to one drug but not to another drug. Response can also refer to the likelihood of successful treatment or the assessment of progress in treatment. Titration of drug therapy to a particular patient is also included in this description, e.g. different patients can respond to different doses of a given medication. This aspect may be important when drugs with side-effects or interactions with other drug therapies are contemplated.

[0207] Diagnostic nucleotide sets are developed and validated for use in assessing whether a patient will respond to a particular therapy and/or monitoring response of a patient to drug therapy(therapies). Disease criteria correspond to presence or absence of clinical symptoms or clinical endpoints, presence of side-effects or interaction with other drug (s). The diagnostic nucleotide set may further comprise nucleotide sequences that are targets of drug treatment or markers of active disease.

Validation and accuracy of diagnostic nucleotide sets

[0208] Prior to widespread application of the diagnostic probe sets of the disclosure the predictive value of the probe set is validated. When the diagnostic probe set is discovered by microarray based expression analysis, the differential expression of the member genes may be validated by a less variable and more quantitive and accurate technology such as real time PCR. In this type of experiment the amplification product is measured during the PCR reaction. This enables the researcher to observe the amplification before any reagent becomes rate limiting for amplification. In kinetic PCR the measurement is of C_T (threshold cycle) or C_P (crossing point). This measurement ($C_T = C_P$) is the point at which an amplification curve crosses a threshold fluorescence value. The threshold is set to a point within the area where all of the reactions were in their linear phase of amplification. When measuring C_T , a lower C_T value is indicative of a higher amount of starting material since an earlier cycle number means the threshold was crossed more quickly.

[0209] Several fluorescence methodologies are available to measure amplification product in real-time PCR. Taqman (Applied BioSystems, Foster City, CA) uses fluorescence resonance energy transfer (FRET) to inhibit signal from a probe until the probe is degraded by the sequence specific binding and Taq 3' exonuclease activity. Molecular Beacons (Stratagene, La Jolla, CA) also use FRET technology, whereby the fluorescence is measured when a hairpin structure is relaxed by the specific probe binding to the amplified DNA. The third commonly used chemistry is Sybr Green, a DNA-binding dye (Molecular Probes, Eugene, OR). The more amplified product that is produced, the higher the signal. The Sybr Green method is sensitive to non-specific amplification products, increasing the importance of primer design

and selection. Other detection chemistries can also been used, such as ethedium bromide or other DNA-binding dyes and many modifications of the fluorescent dye/quencher dye Taqman chemistry, for example scorpions.

[0210] Real-time PCR validation can be done as described in Example 12.

[0211] Typically, the oligonucleotide sequence of each probe is confirmed, e.g. by DNA sequencing using an oligonucleotide-specific primer. Partial sequence obtained is generally sufficient to confirm the identity of the oligonucleotide probe. Alternatively, a complementary polynucleotide is fluorescently labeled and hybridized to the array, or to a different array containing a resynthesized version of the oligo nucleotide probe, and detection of the correct probe is confirmed. [0212] Typically, validation is performed by statistically evaluating the accuracy of the correspondence between the molecular signature for a diagnostic probe set and a selected indicator. For example, the expression differential for a nucleotide sequence between two subject classes can be expressed as a simple ratio of relative expression. The expression of the nucleotide sequence in subjects with selected indicator can be compared to the expression of that nucleotide sequence in subjects without the indicator, as described in the following equations.

the average expression of nucleotide sequence x in the members of group A; $\sum E_x ai/N = E_x A$ the average expression of nucleotide sequence x in the members of group B; $\sum E_x bi/M = E_x B$ the average differential expression of nucleotide sequence x between groups A and B: $E_xA/ExB = \Delta E_xAB$

where Σ indicates a sum; Ex is the expression of nucleotide sequence x relative to a standard; ai are the individual members of group A, group A has N members; bi are the individual members of group B, group B has M members.

[0213] The expression of at least two nucleotide sequences, e.g., nucleotide sequence X and nucleotide sequence Y are measured relative to a standard in at least one subject of group A (e.g., with a disease) and group B (e.g., without the disease). Ideally, for purposes of validation the indicator is independent from (i.e., not assigned based upon) the expression pattern. Alternatively, a minimum threshold of gene expression for nucleotide sequences X and Y, relative to the standard, are designated for assignment to group A. For nucleotide sequence x, this threshold is designated ΔEx , and for nucleotide sequence y, the threshold is designated ΔEy .

[0214] The following formulas are used in the calculations below:

Sensitivity = (true positives/true positives + false negatives)

Specificity = (true negatives/true negatives + false positives)

[0215] If, for example, expression of nucleotide sequence x above a threshold: x > \(\Delta Ex, \) is observed for 80/100 subjects in group A and for 10/100 subjects in group B, the sensitivity of nucleotide sequence x for the assignment to group A, at the given expression threshold ΔEx , is 80%, and the specificity is 90%.

[0216] If the expression of nucleotide sequence y is $> \Delta Ey$ in 80/100 subjects in group A, and in 10/100 subjects in group B, then, similarly the sensitivity of nucleotide sequence y for the assignment to group A at the given threshold Δ Ey is 80% and the specificity is 90%. If in addition, 60 of the 80 subjects in group A that meet the expression threshold for nucleotide sequence y also meet the expression threshold ∆Ex and that 5 of the 10 subjects in group B that meet the expression threshold for nucleotide sequence y also meet the expression threshold ΔEx , the sensitivity of the test (x>\Delta and y>\Delta Ey) for assignment of subjects to group A is 60% and the specificity is 95%.

[0217] Alternatively, if the criteria for assignment to group A are change to: Expression of $x > \Delta Ex$ or expression of y $> \Delta Ey$, the sensitivity approaches 100% and the specificity is 85%.

[0218] Clearly, the predictive accuracy of any diagnostic probe set is dependent on the minimum expression threshold selected. The expression of nucleotide sequence X (relative to a standard) is measured in subjects of groups A (with disease) and B (without disease). The minimum threshold of nucleotide sequence expression for x, required for assignment to group A is designated $\Delta Ex 1$.

[0219] If 90/100 patients in group A have expression of nucleotide sequence $x > \Delta Ex I$ and 20/100 patients in group B have expression of nucleotide sequence $x > \Delta Ex$ I, then the sensitivity of the expression of nucleotide sequence x(using $\Delta Ex\ I$ as a minimum expression threshold) for assignment of patients to group A will be 90% and the specificity will be 80%.

[0220] Altering the minimum expression threshold results in an alteration in the specificity and sensitivity of the nucleotide sequences in question. For example, if the minimum expression threshold of nucleotide sequence x for assignment

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of subjects to group A is lowered to Δ Ex 2, such that 100/100 subjects in group A and 40/100 subjects in group B meet the threshold, then the sensitivity of the test for assignment of subjects to group A will be 100% and the specificity will be 60%.

[0221] Thus, for 2 nucleotide sequences X and Y: the expression of nucleotide sequence x and nucleotide sequence y (relative to a standard) are measured in subjects belonging to groups A (with disease) and B (without disease). Minimum thresholds of nucleotide sequence expression for nucleotide sequences X and Y (relative to common standards) are designated for assignment to group A. For nucleotide sequence x, this threshold is designated Δ Ex1 and for nucleotide sequence y, this threshold is designated Δ Ey1.

[0222] If in group A, 90/100 patients meet the minimum requirements of expression Δ Ex1 and Δ Ey1, and in group B, 10/100 subjects meet the minimum requirements of expression Δ Ex1 and Δ Ey1, then the sensitivity of the test for assignment of subjects to group A is 90% and the specificity is 90%.

[0223] Increasing the minimum expression thresholds for X and Y to Δ Ex2 and Δ Ey2, such that in group A, 70/100 subjects meet the minimum requirements of expression Δ Ex2 and Δ Ey2, and in group B, 3/100 subjects meet the minimum requirements of expression Δ Ex2 and Δ Ey2. Now the sensitivity of the test for assignment of subjects to group A is 70% and the specificity is 97%.

[0224] If the criteria for assignment to group A is that the subject in question meets either threshold, Δ Ex2 or Δ Ey2, and it is found that 100/100 subjects in group A meet the criteria and 20/100 subjects in group B meet the criteria, then the sensitivity of the test for assignment to group A is 100% and the specificity is 80%.

[0225] Individual components of a diagnostic probe set each have a defined sensitivity and specificity for distinguishing between subject groups. Such individual nucleotide sequences can be employed in concert as a diagnostic probe set to increase the sensitivity and specificity of the evaluation. The database of molecular signatures is queried by algorithms to identify the set of nucleotide sequences (i.e., corresponding to members of the probe set) with the highest average differential expression between subject groups. Typically, as the number of nucleotide sequences in the diagnostic probe set increases, so does the predictive value, that is, the sensitivity and specificity of the probe set. When the probe sets are defined they may be used for diagnosis and patient monitoring as discussed below. The diagnostic sensitivity and specificity of the probe sets for the defined use can be determined for a given probe set with specified expression levels as demonstrated above. By altering the expression threshold required for the use of each nucleotide sequence as a diagnostic, the sensitivity and specificity of the probe set can be altered by the practitioner. For example, by lowering the magnitude of the expression differential threshold for each nucleotide sequence in the set, the sensitivity of the test will increase, but the specificity will decrease. As is apparent from the foregoing discussion, sensitivity and specificity are inversely related and the predictive accuracy of the probe set is continuous and dependent on the expression threshold set for each nucleotide sequence. Although sensitivity and specificity tend to have an inverse relationship when expression thresholds are altered, both parameters can be increased as nucleotide sequences with predictive value are added to the diagnostic nucleotide set. In addition a single or a few markers may not be reliable expression markers across a population of patients. This is because of the variability in expression and measurement of expression that exists between measurements, individuals and individuals over time. Inclusion of a large number of candidate nucleotide sequences or large numbers of nucleotide sequences in a diagnostic nucleotide set allows for this variability as not all nucleotide sequences need to meet a threshold for diagnosis. Generally, more markers are better than a single marker. If many markers are used to make a diagnosis, the likelihood that all expression markers will not meet some thresholds based upon random variability is low and thus the test will give fewer false negatives.

[0226] It is appreciated that the desired diagnostic sensitivity and specificity of the diagnostic nucleotide set may vary depending on the intended use of the set. For example, in certain uses, high specificity and high sensitivity are desired. For example, a diagnostic nucleotide set for predicting which patient population may experience side effects may require high sensitivity so as to avoid treating such patients. In other settings, high sensitivity is desired, while reduced specificity may be tolerated. For example, in the case of a beneficial treatment with few side effects, it may be important to identify as many patients as possible (high sensitivity) who will respond to the drug, and treatment of some patients who will not respond is tolerated. In other settings, high specificity is desired and reduced sensitivity may be tolerated. For example, when identifying patients for an early-phase clinical trial, it is important to identify patients who may respond to the particular treatment. Lower sensitivity is tolerated in this setting as it merely results in reduced patients who enroll in the study or requires that more patients are screened for enrollment.

Methods of using diagnostic nucleotide sets.

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[0227] The disclosure also provides methods of using the diagnostic nucleotide sets to: diagnose disease; assess severity of disease; predict future occurrence of disease; predict future complications of disease; determine disease prognosis; evaluate the patient's risk, or "stratify" a group of patients; assess response to current drug therapy; assess response to current non-pharmacological therapy; determine the most appropriate medication or treatment for the patient; predict whether a patient is likely to respond to a particular drug; and determine most appropriate additional diagnostic

testing for the patient, among other clinically and epidemiologically relevant applications.

[0228] The nucleotide sets of the disclosure can be utilized for a variety of purposes by physicians, healthcare workers, hospitals, laboratories, patients, companies and other institutions. As indicated previously, essentially any disease, condition, or status for which at least one nucleotide sequence is differentially expressed in leukocyte populations (or sub-populations) can be evaluated, e.g., diagnosed, monitored, etc. using the diagnostic nucleotide sets and methods of the disclosure. In addition to assessing health status at an individual level, the diagnostic nucleotide sets of the present invention are suitable for evaluating subjects at a "population level," e.g., for epidemiological studies, or for population screening for a condition or disease.

Collection and preparation of sample

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[0229] RNA, protein and/or DNA is prepared using methods well-known in the art, as further described herein. It is appreciated that subject samples collected for use in the methods of the invention are generally collected in a clinical setting, where delays may be introduced before RNA samples are prepared from the subject samples of whole blood, e.g. the blood sample may not be promptly delivered to the clinical lab for further processing. Further delay may be introduced in the clinical lab setting where multiple samples are generally being processed at any given time. For this reason, methods which feature lengthy incubations of intact leukocytes at room temperature are not preferred, because the expression profile of the leukocytes may change during this extended time period. For example, RNA can be isolated from whole blood using a phenol/guanidine isothiocyanate reagent or another direct whole-blood lysis method, as described in, e.g., U.S. Patent Nos. 5,346,994 and 4,843,155. This method may be less preferred under certain circumstances because the large majority of the RNA recovered from whole blood RNA extraction comes from erythrocytes since these cells outnumber leukocytes 1000:1. Care must be taken to ensure that the presence of erythrocyte RNA and protein does not introduce bias in the RNA expression profile data or lead to inadequate sensitivity or specificity of probes.

[0230] Alternatively, intact leukocytes may be collected from whole blood using a lysis buffer that selectively lyses erythrocytes, but not leukocytes, as described, e.g., in (U.S. Patent Nos. 5,973,137, and 6,020,186). Intact leukocytes are then collected by centrifugation, and leukocyte RNA is isolated using standard protocols, as described herein. However, this method does not allow isolation of sub-populations of leukocytes, e.g. mononuclear cells, which may be desired. In addition, the expression profile may change during the lengthy incubation in lysis buffer, especially in a busy clinical lab where large numbers of samples are being prepared at any given time.

[0231] Alternatively, specific leukocyte cell types can be separated using density gradient reagents (Boyum, A, 1968.). For example, mononuclear cells may be separated from whole blood using density gradient centrifugation, as described, e.g., in U.S. Patents Nos. 4190535, 4350593, 4751001, 4818418, and 5053134. Blood is drawn directly into a tube containing an anticoagulant and a density reagent (such as Ficoll or Percoll). Centrifugation of this tube results in separation of blood into an erythrocyte and granulocyte layer, a mononuclear cell suspension, and a plasma layer. The mononuclear cell layer is easily removed and the cells can be collected by centrifugation, lysed, and frozen. Frozen samples are stable until RNA can be isolated. Density centrifugation, however, must be conducted at room temperature, and if processing is unduly lengthy, such as in a busy clinical lab, the expression profile may change.

[0232] Alternatively, cells can be separated using fluorescence activated cell sorting (FACS) or some other technique, which divides cells into subsets based on gene or protein expression. This may be desirable to enrich the sample for cells of interest, but it may also introduce cell manipulations and time delays, which result in alteration of gene expression profiles (Cantor et al. 1975; Galbraith et al. 1999).

[0233] The quality and quantity of each clinical RNA sample is desirably checked before amplification and labeling for array hybridization, using methods known in the art. For example, one microliter of each sample may be analyzed on a Bioanalyzer (Agilent 2100 Palo Alto, CA. USA) using an RNA 6000 nano LabChip (Caliper, Mountain View, CA. USA). Degraded RNA is identified by the reduction of the 28S to 18S ribosomal RNA ratio and/or the presence of large quantities of RNA in the 25-100 nucleotide range.

[0234] It is appreciated that the RNA sample for use with a diagnostic nucleotide set may be produced from the same or a different cell population, sub-population and/or cell type as used to identify the diagnostic nucleotide set. For example, a diagnostic nucleotide set identified using RNA extracted from mononuclear cells may be suitable for analysis of RNA extracted from whole blood or mononuclear cells, depending on the particular characteristics of the members of the diagnostic nucleotide set. Generally, diagnostic nucleotide sets must be tested and validated when used with RNA derived from a different cell population, sub-population or cell type than that used when obtaining the diagnostic gene set. Factors such as the cell-specific gene expression of diagnostic nucleotide set members, redundancy of the information provided by members of the diagnostic nucleotide set, expression level of the member of the diagnostic nucleotide set, and cell-specific alteration of expression of a member of the diagnostic nucleotide set will contribute to the usefullness of using a different RNA source than that used when identifying the members of the diagnostic nucleotide set. It is appreciated that it may be desirable to assay RNA derived from whole blood, obviating the need to isolate particular cell

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Rapid method of RNA extraction suitable for production in a clinical setting of high quality RNA for expression profiling

[0235] In a clinical setting, obtaining high quality RNA preparations suitable for expression profiling, from a desired population of leukocytes poses certain technical challenges, including: the lack of capacity for rapid, high-throughput sample processing in the clinical setting, and the possibility that delay in processing (in a busy lab or in the clinical setting) may adversely affect RNA quality, e.g. by a permitting the expression profile of certain nucleotide sequences to shift. Also, use of toxic and expensive reagents, such as phenol, may be disfavored in the clinical setting due to the added expense associated with shipping and handling such reagents.

[0236] A useful method for RNA isolation for leukocyte expression profiling would allow the isolation of monocyte and lymphocyte RNA in a timely manner, while preserving the expression profiles of the cells, and allowing inexpensive production of reproducible high-quality RNA samples. Accordingly, the invention provides a method of adding inhibitor (s) of RNA transcription and/or inhibitor(s) of protein synthesis, such that the expression profile is "frozen" and RNA degradation is reduced. A desired leukocyte population or sub-population is then isolated, and the sample may be frozen or lysed before further processing to extract the RNA. Blood is drawn from subject population and exposed to ActinontycinD (to a final concentration of 10 ug/ml) to inhibit transcription, and cycloheximide (to a final concentration of 10 ug/ml) to inhibit protein synthesis. The inhibitor(s) can be injected into the blood collection tube in liquid form as soon as the blood is drawn, or the tube can be manufactured to contain either lyophilized inhibitors or inhibitors that are in solution with the anticoagulant. At this point, the blood sample can be stored at room temperature until the desired leukocyte population or sub-population is isolated, as described elsewhere. RNA is isolated using standard methods, e.g., as described above, or a cell pellet or extract can be frozen until further processing of RNA inconvenient.

[0237] Also disclosed is a method of using a low-temperature density gradient for separation of a desired leukocyte sample as well as the combination of use of a low-temperature density gradient and the use of transcriptional and/or protein synthesis inhibitor(s). A desired leukocyte population is separated using a density gradient solution for cell separation that maintains the required density and viscosity for cell separation at 0-4°C. Blood is drawn into a tube containing this solution and may be refrigerated before and during processing as the low temperatures slow cellular processes and minimize expression profile changes. Leukocytes are separated, and RNA is isolated using standard methods. Alternately, a cell pellet or extract is frozen until further processing of RNA is convenient. Care must be taken to avoid rewarming the sample during further processing steps.

[0238] Alternatively, a method of using low-tentperature density gradient separation, combined with the use of actinomycin A and cyclohexamide, as described above can be used.

Assessing expression for diagnostic

[0239] Expression profiles for the set of diagnostic nucleotide sequences in a subject sample can be evaluated by any technique that determines the expression of each component nucleotide sequence. Methods suitable for expression analysis are known in the art, and numerous examples are discussed in the Sections titled "Methods of obtaining expression data" and "high throughput expression Assays", above.

[0240] In many cases, evaluation of expression profiles is most efficiently, and cost effectively, performed by analyzing RNA expression. Alternatively, the proteins encoded by each component of the diagnostic nucleotide set are detected for diagnostic purposes by any technique capable of determining protein expression, e.g., as described above. Expression profiles can be assessed in subject leukocyte sample using the same or different techniques as those used to identify and validate the diagnostic nucleotide set, For example, a diagnostic nucleotide set identified as a subset of sequences on a cDNA microarray can be utilized for diagnostic (or prognostic, or monitoring, etc.) purposes on the same array from which they were identified. Alternatively, the diagnostic nucleotide sets for a given disease or condition can be organized onto a dedicated sub-array for the indicated purpose. It is important to note that if diagnostic nucleotide sets are discovered using one technology, e.g. RNA expression profiling, but applied as a diagnostic using another technology, e.g. protein expression profiling, the nucleotide sets must generally be validated for diagnostic purposes with the new technology. In addition, it is appreciated that diagnostic nucleotide sets that are developed for one use, e.g. to diagnose a particular disease, may later be found to be useful for a different application, e.g. to predict the likelihood that the particular disease will occur. Generally, the diagnostic nucleotide set will need to be validated for use in the second circumstance. As discussed herein, the sequence of diagnostic nucleotide set members may be amplified from RNA or cDNA using methods known in the art providing specific amplification of the nucleotide sequences.

General Protein Methods

[0241] Protein products of the nucleotide sequences disclosed herein may include proteins that represent functionally

equivalent gene products. Such an equivalent gene product may contain deletions, additions or substitutions of amino acid residues within the amino acid sequence encoded by the nucleotide sequences described, above, but which result in a silent change, thus producing a functionally equivalent nucleotide sequence product. Amino acid, substitutions may be made on the basis of similarity in polarity, charge, solubility, hydrophobicity, hydrophilicity, and/or the amphipathic nature of the residues involved.

[0242] For example, nonpolar (hydrophobic) amino acids include alanine, leucine, isoleucine, valine, proline, phenylalanine, tryptophan, and methionine; polar neutral amino acids include glycine, serine, threonine, cysteine, tyrosine, asparagine, and glutamine; positively charged (basic) amino acids include arginine, lysine, and histidine; and negatively charged (acidic) amino acids include aspartic acid and glutamic acid. "Functionally equivalent", as utilized herein, refers to a protein capable of exhibiting a substantially similar in vivo activity as the endogenous gene products encoded by the nucleotide described, above.

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[0243] The gene products (protein products of the nucleotide sequences) may be produced by recombinant DNA technology using techniques well known in the art. Thus, methods for preparing the gene polypeptides and peptides of the invention by expressing nucleic acid encoding nucleotide sequences are described herein. Methods which are well known to those skilled in the art can be used to construct expression vectors containing nucleotide sequence protein coding sequences and appropriate transcriptional/translational control signals. These methods include, for example, in vitro recombinant DNA techniques, synthetic techniques and in vivo recombination/genetic recombination. See, for example, the techniques described in Sambrook et al., 1989, supra, and Ausubel et al., 1989, supra. Alternatively, RNA capable of encoding nucleotide sequence protein sequences may be chemically synthesized using, for example, synthesizers. See, for example, the techniques described in "Oligonucleotide Synthesis", 1984, Gait, M. J. ed., IRL Press, Oxford.

[0244] A variety of host-expression vector systems may be utilized to express the nucleotide sequence coding sequences of the disclosure. Such host-expression systems represent vehicles by which the coding sequences of interest may be produced and subsequently purified, but also represent cells which may, when transformed or transfected with the appropriate nucleotide coding sequences, exhibit the protein encoded by the nucleotide sequence of the invention in situ. These include but are not limited to microorganisms such as bacteria (e.g., E. coli, B. subtilis) transformed with recombinant bacteriophage DNA, plasmid DNA or cosmid DNA expression vectors containing nucleotide sequence protein coding sequences; yeast (e.g. Saccharomyces, Pichia) transformed with recombinant yeast expression vectors containing the nucleotide sequence protein coding sequences; insect cell systems infected with recombinant virus expression vectors (e.g., baculovirus) containing the nucleotide sequence protein coding sequences; plant cell systems infected with recombinant virus expression vectors (e.g., cauliflower mosaic virus, CaMV; tobacco mosaic virus, TMV) or transformed with recombinant plasmid expression vectors (e.g., Ti plasmid) containing nucleotide sequence protein coding sequences; or mammalian cell systems (e.g. COS, CHO, BHK, 293, 3T3) harboring recombinant expression constructs containing promoters derived from the genome of mammalian cells (e.g., metallothionein promoter) or from mammalian viruses (e.g., the adenovirus late promoter; the vaccinia virus 7.5 K promoter).

[0245] In bacterial systems, a number of expression vectors may be advantageously selected depending upon the use intended for the nucleotide sequence protein being expressed. For example, when a large quantity of such a protein is to be produced, for the generation of antibodies or to screen peptide libraries, for example, vectors which direct the expression of high levels of fusion protein products that are readily purified may be desirable. Such vectors include, but are not limited, to the E. coli expression vector pUR278 (Ruther et al., 1983, EMBO J. 2:1791), in which the nucleotide sequence protein coding sequence may be ligated individually into the vector in frame with the lac Z coding region so that a fusion protein is produced; pIN vectors (Inouye & Inouye, 1985, Nucleic Acids Res. 13:3101-3109; Van Heeke & Schuster, 1989, J. Biol. Chem. 264:5503-5509); and the likes of pGEX vectors may also be used to express foreign polypeptides as fusion proteins with glutathione S-transferase (GST). In general, such fusion proteins are soluble and can easily be purified from lysed cells by adsorption to glutathione-agarose beads followed by elution in the presence of free glutathione. The pGEX vectors are designed to include thrombin or factor Xa protease cleavage sites so that the cloned target nucleotide sequence protein can be released from the GST moiety. Other systems useful in the disclosure include use of the FLAG epitope or the 6-HIS systems.

[0246] In an insect system, Autographa californica nuclear polyhedrosis virus (AcNPV) is used as a vector to express foreign nucleotide sequences. The virus grows in Spodoptera frugiperda cells. The nucleotide sequence coding sequence may be cloned individually into non-essential regions (for example the polyhedrin gene) of the virus and placed under control of an AcNPV promoter (for example the polyhedrin promoter). Successful insertion of nucleotide sequence coding sequence will result in inactivation of the polyhedrin gene and production of non-occluded recombinant virus (i.e., virus lacking the proteinaceous coat coded for by the polyhedrin gene). These recombinant viruses are then used to infect Spodoptera frugiperda cells in which the inserted nucleotide sequence is expressed. (E.g., see Smith et al., 1983, J. Virol. 46: 584; Smith, U.S. Pat. No. 4,215,051).

[0247] In mammalian host cells, a number of viral-based expression systems may be utilized. In cases where an adenovirus is used as an expression vector, the nucleotide sequence coding sequence of interest may be ligated to an

adenovirus transcription/translation control complex, e.g., the late promoter and tripartite leader sequence. This chimeric nucleotide sequence may then be inserted in the adenovirus genome by in vitro or in vivo recombination. Insertion in a non-essential region of the viral genome (e.g., region E1 or E3) will result in a recombinant virus that is viable and capable of expressing nucleotide sequence encoded protein in infected hosts. (E.g., See Logan & Shenk, 1984, Proc. Natl. Acad. Sci. USA 81:3655-3659). Specific initiation signals may also be required for efficient translation of inserted nucleotide sequence coding sequences. These signals include the ATG initiation codon and adjacent sequences. In cases where an entire nucleotide sequence, including its own initiation codon and adjacent sequences, is inserted into the appropriate expression vector, no additional translational control signals may be needed. However, in cases where only a portion of the nucleotide sequence coding sequence is inserted, exogenous translational control signals, including, perhaps, the ATG initiation codon, must be provided. Furthermore, the initiation codon must be in phase with the reading frame of the desired coding sequence to ensure translation of the entire insert. These exogenous translational control signals and initiation codons can be of a variety of origins, both natural and synthetic. The efficiency of expression may be enhanced by the inclusion of appropriate transcription enhancer elements, transcription terminators, etc. (see Bittner et al., 1987, Methods in Enzymol. 153:516-544).

[0248] In addition, a host cell strain may be chosen which modulates the expression of the inserted sequences, or modifies and processes the product of the nucleotide sequence in the specific fashion desired. Such modifications (e.g., glycosylation) and processing (e.g., cleavage) of protein products may be important for the function of the protein. Different host cells have characteristic and specific mechanisms for the post-translational processing and modification of proteins. Appropriate cell lines or host systems can be chosen to ensure the correct modification and processing of the foreign protein expressed. To this end, eukaryotic host cells which possess the cellular machinery for proper processing of the primary transcript, glycosylation, and phosphorylation of the gene product may be used. Such mammalian host cells include but are not limited to CHO, VERO, BHK, HeLa, COS, MDCK, 293, 3T3, WI38, etc.

[0249] For long-term, high-yield production of recombinant proteins, stable expression is preferred. For example, cell lines which stably express the nucleotide sequence encoded protein may be engineered. Rather than using expression vectors which contain viral origins of replication, host cells can be transformed with DNA controlled by appropriate expression control elements (e.g., promoter, enhancer, sequences, inscription terminators, polyadenylation sites, etc.), and a selectable marker. Following the introduction of the foreign DNA, engineered cells may be allowed to grow for 1-2 days in an enriched media, and then are switched to a selective media. The selectable marker in the recombinant plasmid confers resistance to the selection and allows cells to stably integrate the plasmid into their chromosome and grow to form foci which in turn can be cloned and expanded into cell lines. This method may advantageously be used to engineer cell lines which express nucleotide sequence encoded protein. Such engineered cell lines may be particularly useful in screening and evaluation of compounds that affect the endogenous activity of the nucleotide sequence encoded protein. [0250] A number of selection systems may be used, including but not limited to the herpes simplex virus thymidine kinase (Wigler, et al., 1977, Cell 11:223), hypoxanthine-guanine phosphoribosyltransferase (Szybalska & Szybalski, 1962, Proc. Natl. Acad. Sci. USA 48:2026), and adenime phosphoribosyltransferase (Lowy, et al., 1980, Cell 22:817) genes can be employed in tk-, hgprt- or aprt- cells, respectively. Also, antimetabolite resistance can be used as the basis of selection for dhfr, which confers resistance to methotrexate (Wigler, et al., 1980, Natl. Acad. Sci. USA 77:3567; O'Hare, et al., 1981, Proc. NatL Acad. Sci. USA 78:1527); gpt, which confers resistance to mycophenolic acid (Mulligan & Berg, 1981, Proc. Natl. Acad. Sci. USA 78:2072); neo, which confers resistance to the aminoglycoside G-418 (Colberre-Garapin, et al., 1981, J. Mol. Biol. 150:1); and hygro, which confers resistance to hygromycin (Santerre, et al., 1984, Gene 30:147) genes.

[0251] An alternative fusion protein system allows for the ready purification of non-danatured fusion proteins expressed in human cell lines (Janknecht, et al., 1991, Proc. Natl. Acad. Sci. USA 88: 8972-8976). In this system, the nucleotide sequence of interest is subcloned into a vaccinia recombination plasmid such that the nucleotide sequence's open reading frame is translationally fused to an amino-terminal tag consisting of six histidine residues. Extracts from cells infected with recombinant vaccinia virus are loaded onto Ni.sup.2 +-nitriloacetic acid-agarose columns and histidine-tagged proteins are selectively eluted with imidazole-containing buffers.

[0252] Where recombinant DNA technology is used to produce the protein encoded by the nucleotide sequence for such assay systems, it may be advantageous to engineer fusion proteins that can facilitate labeling, immobilization and/or detection.

Antibodies

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[0253] Indirect labeling involves the use of a protein, such as a labeled antibody, which specifically binds to the protein encoded by the nucleotide sequence. Such antibodies include but are not limited to polyclonal, monoclonal, chimeric, single chain, Fab fragments and fragments produced by an Fab expression library.

[0254] Antibodies to the protein encoded by the nucleotide sequences are also disclosed. Described herein are methods for the production of antibodies capable of specifically recognizing one or more nucleotide sequence epitopes. Such

antibodies may include, but are not limited to polyclonal antibodies, monoclonal antibodies (mAbs), humanized or chimeric antibodies, single chain antibodies, Fab fragments, F(ab')2 fragments, fragments produced by a Pub expression library, anti-idiotypic (anti-Id) antibodies, and epitope-binding fragments of any of the above. Such antibodies may be used, for example, in the detection of a nucleotide sequence in a biological sample, or, alternatively, as a method for the inhibition of abnormal gene activity, for example, the inhibition of a disease target nucleotide sequence, as further described below. Thus, such antibodies may be utilized as part of cardiovascular or other disease treatment method, and/or may be used as part of diagnostic techniques whereby patients may be tested for abnormal levels of nucleotide sequence encoded proteins, or for the presence of abnormal forms of the such proteins.

[0255] For the production of antibodies to a nucleotide sequence, various host animals may be immunized by injection with a protein encoded by the nucleotide sequence, or a portion thereof. Such host animals may include but are not limited to rabbits, mice, and rats, to name but a few. Various adjuvants may be used to increase the immunological response, depending on the host species, including but not limited to Freund's (complete and incomplete), mineral gels such as aluminum hydroxide, surface active substances such as lysolecithin, pluronic polyols, polyanions, peptides, oil emulsions, keyhole limpet hemocyanin, dinitrophenol, and potentially useful human adjuvants such as BCG (bacille Calmette-Guerin) and Corynebacterium parvum.

[0256] Polyclonal antibodies are heterogeneous populations of antibody molecules derived from the sera of animals immunized with an antigen, such as gene product, or an antigenic functional derivative thereof. For the production of polyclonal antibodies, host animals such as those described above, may be immunized by injection with gene product supplemented with adjuvants as also described above.

[0257] Monoclonal antibodies, which are homogeneous populations of antibodies to a particular antigen, may be obtained by any technique which provides for the production of antibody molecules by continuous cell lines in culture. These include, but are not limited to the hybridoma technique of Kohler and Milstein, (1975, Nature 256:495-497; and U.S. Pat. No. 4,376,110), the human B-cell hybridoma technique (Kosbor et al., 1983, Immunology Today 4:72; Cole et al., 1983, Proc. Natl. Acad. Sci. USA 80:2026-2030), and the EBV-hybridoma technique (Cole et al., 1985, Monoclonal Antibodies And Cancer Therapy, Alan R. Liss, Inc., pp. 77-96). Such antibodies may be of any immunoglobulin class including IgG, IgM, IgE, IgA, IgD and any subclass thereof. The hybridoma producing the mAb of this invention may be cultivated in vitro or in vivo.

[0258] In addition, techniques developed for the production of "chimeric antibodies" (Morrison et al., 1984, Proc. Natl. Acad. Sci., 81:6851-6855; Neuberger et al., 1984, Nature, 312:604-608; Takeda et al., 1985, Nature, 314:452-454) by splicing the genes from a mouse antibody molecule of appropriate antigen specificity together with genes from a human antibody molecule of appropriate biological activity can be used. A chimeric antibody is a molecule in which different portions are derived from different animal species, such as those having a variable region derived from a murine mAb and a human immunoglobulin constant region.

[0259] Alternatively, techniques described for the production of single chain antibodies (U.S. Pat. No. 4,946,778; Bird, 1988, Science 242:423-426; Huston et al., 1988, Proc. Natl. Acad. Sci. USA 85:5879-5883; and Ward et al., 1989, Nature 334:544-546) can be adapted to produce nucleotide sequence-single chain antibodies. Single chain antibodies are formed by linking the heavy and light chain fragments of the Fv region via an amino acid bridge, resulting in a single chain polypeptide.

[0260] Antibody fragments which recognize specific epitopes may be generated by known techniques For example, such fragments include but are not limited to the F(ab')2 fragments which can be produced by pepsin digestion of the antibody molecule and the Fab fragments which can be generated by reducing the disulfide bridges of the F(ab')2 fragments. Alternatively, Fab expression libraries may be constructed (Huse et al., 1989, Science, 246:1275-1281) to allow rapid and easy identification of Monoclonal Fab fragments with the desired specificity.

45 <u>Disease specific target nucleotide sequences</u>

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[0261] Disease specific target nucleotide sequences, and sets of disease specific target nucleotide sequences are also disclosed. The diagnostics nucleotide sets, subsets thereof, novel nucleotide sequences, and individual members of the diagnostic nucleotide sets identified as described above are also disease specific target nucleotide sequences. In particular, individual nucleotide sequences that are differentially regulated or have predictive value that is strongly correlated with a disease or disease criterion are especially favorable as disease specific target nucleotide sequences. Sets of genes that are co-regulated may also be identified as disease specific target nucleotide sets. Such nucleotide sequences and/or nucleotide sequence products are targets for modulation by a variety of agents and techniques. For example, disease specific target nucleotide sequences (or the products of such nucleotide sequences, or sets of disease specific target nucleotide sequences) can be inhibited or activated by, e.g., target specific monoclonal antibodies or small molecule inhibitors, or delivery of the nucleotide sequence or gene product of the nucleotide sequence to patients. Also, sets of genes can be inhibited or activated by a variety of agents and techniques. The specific usefulness of the target nucleotide sequence(s) depends on the subject groups from which they were discovered, and the disease or

disease criterion with which they correlate.

Imaging

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Further disclosed are imaging reagents. The differentially expressed leukocyte nucleotide sequences, diagnostic nucleotide sets, or portions thereof, and novel nucleotide sequences of the invention are nucleotide sequences expressed in cells with or without disease. Leukocytes expressing a nucleotide sequence(s) that is differentially expressed in a disease condition may localize within the body to sites that are of interest for imaging purposes. For example, a leukocyte expressing a nucleotide sequence(s) that are differentially expressed in an individual having atherosclerosis may localize or accumulate at the site of an atherosclerotic placque. Such leukocytes, when labeled, may provide a detection reagent for use in imaging regions of the body where labeled leukocyte accumulate or localize, for example, at the atherosclerotic plaque in the case of atherosclerosis. For example, leukocytes are collected from a subject, labeled in vitro, and reintroduced into a subject. Alternatively, the labeled reagent is introduced into the subject individual, and leukocyte labeling occurs within the patient.

[0263] Imaging agents that detect the imaging targets of the invention are produced by well-known molecular and immunological methods (for exemplary protocols, see, e.g., Ausubel, Berger, and Sambrook, as well as Harlow and Lane, supra).

[0264] For example, a full-length nucleic acid sequence, or alternatively, a gene fragment encoding an immunogenic peptide or polypeptide fragment, is cloned into a convenient expression vector, for example, a vector including an inframe epitope or substrate binding tag to facilitate subsequent purification. Protein is then expressed from the cloned cDNA sequence and used to generate antibodies, or other specific binding molecules, to one or more antigens of the imaging target protein. Alternatively, a natural or synthetic polypeptide (or peptide) or small molecule that specifically binds (or is specifically bound to) the expressed imaging target can be identified through well established techniques (see, e.g., Mendel et al, (2006) Anticancer Drug Des 15:29-41; Wilson (2000) Curr Med Chem 7:73-98; Hamby and Showwalter (1999) Pharmacol Ther 82:169-93; and Shimazawa et al. (1998) Curr Opin Struct Biol 8:451-8). The binding molecule, e.g., antibody, small molecule ligand, etc., is labeled with a contrast agent or other detectable label, e.g., gadolinium, iodine, or a gamma-emitting source. For in-vivo imaging of a disease process that involved leukocytes, the labeled antibody is infused into a subject, e.g., a human patient or animal subject, and a sufficient period of time is passed to permit binding of the antibody to target cells. The subject is then imaged with appropriate technology such as MRI (when the label is gadolinium) or with a gamma counter (when the label is a gamma emitter).

<u>Identification of nucleotide sequence involved in adhesion</u>

[0265] A method of identifying nucleotide sequences involved in leukocyte adhesion is described. The interaction between the endothelial cell and leukocyte is a fundamental mechanism of all inflammatory disorders, including the diagnosis and prognosis of allograft rejection the diseases listed in Table 1. For example, the first visible abnormality in atherosclerosis is the adhesion to the endothelium and diapedesis of mononuclear cells (e.g., T-cell and monocyte). Insults to the endothelium (for example, cytokines, tobacco, diabetes, hypertension and many more) lead to endothelial cell activation. The endothelium then expresses adhesion molecules, which have counter receptors on mononuclear cells. Once the leukocyte receptors have bound the endothelial adhesion molecules, they stick to the endothelium, roll a short distance, stop and transmigrate across the endothelium. A similar set of events occurs in both acute and chronic inflammation. When the leukocyte binds the endothelial adhesion molecule, or to soluble cytokines secreted by endothelial or other cells, a program of gene expression is activated in the leukocyte. This program of expression leads to leukocyte rolling, firm adhesion and transmigration into the vessel wall or tissue parenchyma. Inhibition of this process is highly desirable goal in anti-inflammatory drug development. In addition, leukocyte nucleotide sequences and epithelial cell nucleotide sequences, that are differentially expressed during this process may be disease-specific target nucleotide sequences.

[0266] Human endothelial cells, e.g. derived from human coronary arteries, human aorta, human pulmonary artery, human umbilical vein or microvascular endothelial cells, are cultured as a confluent monolayer, using standard methods. Some of the endothelial cells are then exposed to cytokines or another activating stimuli such as oxidized LDL, hypergylycemia, shear stress, or hypoxia (Moser et al. 1992). Some endothelial cells are not exposed to such stimuli and serve as controls. For example, the endothelial cell monolayer is incubated with culture medium containing 5 U/ml of human recombinant IL-1alpha or 10 ng/ml TNF (tumor necrosis factor), for a period of minutes to overnight. The culture medium composition is changed or the flask is sealed to induce hypoxia. In addition, tissue culture plate is rotated to induce sheer stress

[0267] Human T-cells and/or monocytes are cultured in tissue culture flasks or plates, with LGM-3 media from Clonetics. Cells are incubated at 37 degree C, 5% CO2 and 95% humidity. These leukocytes are exposed to the activated or control endothelial layer by adding a suspension of leukocytes on to the endothelial cell monolayer. The endothelial cell monolayer

is cultured on a tissue culture treated plate/ flask or on a microporous membrane. After a variable duration of exposures, the endothelial cells and leukocytes are harvested separately by treating all cells with trypsin and then sorting the endothelial cells from the leukocytes by magnetic affinity reagents to an endothelial cell specific marker such as PECAM-1 (Stem Cell Technologies). RNA is extracted from the isolated cells by standard techniques. Leukocyte RNA is labeled as described above, and hybridized to leukocyte candidate nucleotide library. Epithelial cell RNA is also labeled and hybridized to the leukocyte candidate nucleotide library. Alternatively, the epithelial cell RNA is hybridized to a epithelial cell candidate nucleotide library, prepared according to the methods described for leukocyte candidate libraries, above. [0268] Hybridization to candidate nucleotide libraries will reveal nucleotide sequences that are upregulated or down-regulated in leukocyte and/or epithelial cells undergoing adhesion. The differentially regulated nucleotide sequences are further characterized, e.g. by isolating and sequencing the full-length sequence, analysis of the DNA and predicted protein sequence, and functional characterization of the protein product of the nucleotide sequence, as described above. Further characterization may result in the identification of leukocyte adhesion specific target nucleotide sequences, which may be candidate targets for regulation of the inflammatory process. Small molecule or antibody inhibitors can be developed to inhibit the target nucleotide sequence function. Such inhibitors are tested for their ability to inhibit leukocyte adhesion in the in vitro test described above.

Integrated systems

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[0269] Integrated systems for the collection and analysis of expression profiles, and molecular signatures, as well as for the compilation, storage and access of the databases of the invention, typically include a digital computer with software including an instruction set for sequence searching and analysis, and, optionally, high-throughput liquid control software, image analysis software, data interpretation software, a robotic control armature for transferring solutions from a source to a destination (such as a detection device) operably linked to the digital computer, an input device (e.g., a computer keyboard) for entering subject data to the digital computer, or to control analysis operations or high throughput sample transfer by the robotic control armature. Optionally, the integrated system further comprises an image scanner for digitizing label signals from labeled assay components, e.g., labeled nucleic acid hybridized to a candidate library microarray. The image scanner can interface with image analysis software to provide a measurement of the presence or intensity of the hybridized label, i.e., indicative of an on/off expression pattern or an increase or decrease in expression. [0270] Readily available computational hardware resources using standard operating systems are fully adequate, e.g., a PC (Intel x86 or Pentium chip- compatible DOS,™ OS2,™ WINDOWS,™ WINDOWS NT,™ WINDOWS95,™ WINDOWS98,™ LINUX, or even Macintosh, Sun or PCs will suffice) for use in the integrated systems of the disclosure. Current art in software technology is similarly adequate (i.e., there are a multitude of mature programming languages and source code suppliers) for design, e.g., of an upgradeable open-architecture object-oriented heuristic algorithm, or instruction set for expression analysis, as described herein. For example, software for aligning or otherwise manipulating, molecular signatures can be constructed by one of skill using a standard programming language such as Visual basic, Fortran, Basic, Java, or the like, according to the methods herein.

[0271] Various methods and algorithms, including genetic algorithms and neural networks, can be used to perform the data collection, correlation, and storage functions, as well as other desirable functions, as described herein. In addition, digital or analog systems such as digital or analog computer systems can control a variety of other functions such as the display and/or control of input and output files.

[0272] For example, standard desktop applications such as word processing software (e.g., Corel WordPerfect™ or Microsoft Word™) and database software (e.g., spreadsheet software such as Corel Quattro Pro™, Microsoft Excel™, or database programs such as Microsoft Access™ or Paradox™) can be adapted to the present invention by inputting one or more character string corresponding, e.g., to an expression pattern or profile, subject medical or historical data, molecular signature, or the like, into the software which is loaded into the memory of a digital system, and carrying out the operations indicated in an instruction set, e.g., as exemplified in Figure 2. For example, systems can include the foregoing software having the appropriate character string information, e.g., used in conjunction with a user interface in conjunction with a standard operating system such as a Windows, Macintosh or LINUX system. For example, an instruction set for manipulating strings of characters, either by programming the required operations into the applications or with the required operations performed manually by a user (or both). For example, specialized sequence alignment programs such as PILEUP or BLAST can also be incorporated into the systems, e.g., for alignment of nucleic acids or proteins (or corresponding character strings).

[0273] Software for performing the statistical methods required, e.g., to determine correlations between expression profiles and subsets of members of the diagnostic nucleotide libraries, such as programmed embodiments of the statistical methods described above, are also included in the computer systems. Alternatively, programming elements for performing such methods as principle component analysis (PCA) or least squares analysis can also be included in the digital system to identify relationships between data. Exemplary software for such methods is provided by Partek, Inc., St. Peter, Mo; at the web site partek.com.

[0274] Any controller or computer optionally includes a monitor which can include, e.g., a flat panel display (e.g., active matrix liquid crystal display, liquid crystal display), a cathode ray tube ("CRT") display, or another display system which serves as a user interface, e.g., to output predictive data. Computer circuitry, including numerous integrated circuit chips, such as a microprocessor, memory, interface circuits, and the like, is often placed in a casing or box which optionally also includes a hard disk drive, a floppy disk drive, a high capacity removable drive such as a writeable CD-ROM, and other common peripheral elements.

[0275] Inputting devices such as a keyboard, 'mouse, or touch sensitive screen, optionally provide for input from a user and for user selection, e.g., of sequences or data sets to be compared or otherwise manipulated in the relevant computer system. The computer typically includes appropriate software for receiving user instructions, either in the form of user input into a set parameter or data fields (e.g., to input relevant subject data), or in the form of preprogrammed instructions, e.g., preprogrammed for a variety of different specific operations. The software then converts these instructions to appropriate language for instructing the system to carry out any desired operation.

[0276] The integrated system may also be embodied within the circuitry of an application specific integrated circuit (ASIC) or programmable logic device (PLD). In such a case, the invention is embodied in a computer readable descriptor language that can be used to create an ASIC or PLD. The integrated system can also be embodied within the circuitry or logic processors of a variety of other digital apparatus, such as PDAs, laptop computer systems, displays, image editing equipment, etc.

[0277] The digital system can comprise a learning component where expression profiles, and relevant subject data are compiled and monitored in conjunction with physical assays, and where correlations, e.g., molecular signatures with predictive value for a disease, are established or refined. Successful and unsuccessful combinations are optionally documented in a database to provide justification/preferences for user-base or digital system based selection of diagnostic nucleotide sets with high predictive accuracy for a specified disease or condition.

[0278] The integrated systems can also include an automated workstation. For example, such a workstation can prepare and analyze leukocyte RNA samples by performing a sequence of events including: preparing RNA from a human blood sample; labeling the RNA with an isotopic or non-isotopic label; hybridizing the labeled RNA to at least one array comprising all or part of the candidate library; and detecting the hybridization pattern. The hybridization pattern is digitized and recorded in the appropriate database.

Automated RNA preparation tool

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[0279] An automated RNA preparation tool for the preparation of mononulclear cells from whole blood samples, and preparation of RNA from the mononuclear cells is also disclosed. In a preferred embodiment, the use of the RNA preparation tool is fully automated, so that the cell separation and RNA isolation would require no human manipulations. Full automation is advantageous because it minimizes delay, and standardizes sample preparation across different laboratories. This standardization increases the reproducibility of the results.

[0280] Figure 2 depicts the processes performed by the RNA preparation tool. A primary component of the device is a centrifuge (A). Tubes of whole blood containing a density gradient solution, transcription/translation inhibitors, and a gel barrier that separates erythrocytes from mononuclear cells and serum after centrifugation are placed in the centrifuge (B). The barrier is permeable to erythrocytes and granulocytes during centrifugation, but does not allow mononuclear cells to pass through (or the barrier substance has a density such that mononuclear cells remain above the level of the barrier during the centrifugation). After centrifugation, the erythrocytes and granulocytes are trapped beneath the barrier, facilitating isolation of the mononuclear cell and serum layers. A mechanical arm removes the tube and inverts It to mix the mononuclear cell layer and the serum (C). The arm next pours the supernatant into a fresh tube (D), while the erythrocytes and granulocytes remained below the barrier. Alternatively, a needle is used to aspirate the supernatant and transfer it to a fresh tube. The mechanical arms of the device opens and closes lids, dispenses PBS to aid in the collection of the mononuclear cells by centrifugation, and moves the tubes in and out of the centrifuge. Following centrifugation, the supernatant is poured off or removed by a vacuum device (E), leaving an isolated mononuclear cell pellet. Purification of the RNA from the cells is performed automatically, with lysis buffer and other purification solutions (F) automatically dispensed and removed before and after centrifugation steps. The result is a purified RNA solution. In another embodiment, RNA isolation is performed using a column or filter method. In yet another embodiment, the invention includes an on-board homogenizer for use in cell lysis.

Other automated systems

[0281] Automated and/or semi-automated methods for solid and liquid phase high-throughput sample preparation and evaluation are available, and supported by commercially available devices. For example, robotic devices for preparation of nucleic acids from bacterial colonies, e.g., to facilitate production and characterization of the candidate library include, for example, an automated colony picker (e.g., the Q-bot, Genetix, U.K.) capable of identifying, sampling, and inoculating

up to 10,000/4 hrs different clones into 96 well microtiter dishes. Alternatively, or in addition, robotic systems for liquid handling are available from a variety of sources, e.g., automated workstations like the automated synthesis apparatus developed by Takeda Chemical Industries, LTD. (Osaka, Japan) and many robotic systems utilizing robotic arms (Zymate II, Zymark Corporation, Hopkinton, Mass.; Orca, Beckman Coulter, Inc. (Fullerton, CA)) which mimic the manual operations performed by a scientist Any of the above devices are suitable for use with the present invention, e.g., for high-throughput analysis of library components or subject leukocyte samples. The nature and implementation of modifications to these devices (if any) so that they can operate as discussed herein will be apparent to persons skilled in the relevant art. [0282] High throughput screening systems that automate entire procedures, e.g., sample and reagent pipetting, liquid dispensing, timed incubations, and final readings of the microplate in detector(s) appropriate for the relevant assay are commercially available. (see, e.g., Zymark Corp., Hopkinton, MA; Air Technical Industries, Mentor, OH; Beckman Instruments, Inc. Fullerton, CA; Precision Systems, Inc., Natick, MA, etc.). These configurable systems provide high throughput and rapid start up as well as a high degree of flexibility and customization. Similarly, arrays and array readers are available, e.g., from Affymetrix, PE Biosystems, and others.

[0283] The manufacturers of such systems provide detailed protocols the various high throughput. Thus, for example, Zymark Corp. provides technical bulletins describing screening systems for detecting the modulation of gene transcription, ligand binding, and the like.

[0284] A variety of commercially available peripheral equipment, including, e.g., optical and fluorescent detectors, optical and fluorescent microscopes, plate readers, CCD arrays, phosphorimagers, scintillation counters, phototubes, photodiodes, and the like, and software is available for digitizing, storing and analyzing a digitized video or digitized optical or other assay results, e.g., using PC (Intel x86 or pentium chip- compatible DOS™, OS2™ WINDOWS™, WINDOWS NT™ or WINDOWS95™ based machines), MACINTOSH™, or UNIX based (e.g., SUN™ work station) computers.

Embodiment in a web site.

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[0285] The methods described above can be implemented in a localized or distributed computing environment. For example, if a localized computing environment is used, an array comprising a candidate nucleotide library, or diagnostic nucleotide set, is configured in proximity to a detector, which is, in turn, linked to a computational device equipped with user input and output features.

[0286] In a distributed environment, the methods can be implemented on a single computer with multiple processors or, alternatively, on multiple computers. The computers can be linked, e.g. through a shared bus, but more commonly, the computer(s) are nodes on a network. The network can be generalized or dedicated, at a local level or distributed over a wide geographic area. In certain embodiment, the computers are components of an intra-net or an internet.

[0287] The predictive data corresponding to subject molecular signatures (e.g., expression profiles, and related diagnostic, prognostic, or monitoring results) can be shared by a variety of parties. In particular, such information can be utilized by the subject, the subject's health care practitioner or provider, a company or other institution, or a scientist. An individual subject's data, a subset of the database or the entire database recorded in a computer readable medium can be accessed directly by a user by any method of communication, including, but not limited to, the internet. With appropriate computational devices, integrated systems, communications networks, users at remote locations, as well as users located in proximity to, e.g., at the same physical facility, the database can access the recorded information. Optionally, access to the database can be controlled using unique alphanumeric passwords that provide access to a subset of the data. Such provisions can be used, e.g., to ensure privacy, anonymity, etc.

[0288] Typically, a client (e.g., a patient, practitioner, provider, scientist, or the like) executes a Web browser and is linked to a server computer executing a Web server. The Web browser is, for example, a program such as IBM's Web Explorer, Internet explorer, NetScape or Mosaic, or the like. The Web server is typically, but not necessarily, a program such as IBM's HTTP Daemon or other WWW daemon (e.g., LINUX-based forms of the program). The client computer is bi-directionally coupled with the server computer over a line or via a wireless system. In turn, the server computer is bi-directionally coupled with a website (server hosting the website) providing access to software implementing the methods of this invention.

[0289] A user of a client connected to the Intranet or Internet may cause the client to request resources that are part of the web site(s) hosting the application(s) providing an implementation of the methods described herein. Server program (s) then process the request to return the specified resources (assuming they are currently available). A standard naming convention has been adopted, known as a Uniform Resource Locator ("URL"). This convention encompasses several types of location names, presently including subclasses such as Hypertext Transport Protocol ("http"), File Transport Protocol ("ftp"), gopher, and Wide Area Information Service ("WAIS"). When a resource is downloaded, it may include the URLs of additional resources. Thus, the user of the client can easily learn of the existence of new resources that he or she had not specifically requested.

[0290] Methods of implementing Intranet and/or Intranet embodiments of computational and/or data access processes

are well known to those of skill in the art and are documented, e.g., in ACM Press, pp. 383-392; ISO-ANSI, Working Draft, "Information Technology-Database Language SQL", Jim Melton, Editor, International Organization for Standardization and American National Standards Institute, Jul. 1992; ISO Working Draft, "Database Language SQL-Part 2: Foundation (SQL/Foundation)", CD9075-2:199.chi.SQL, Sep. 11, 1997; and Cluer et al. (1992) A General Framework for the Optimization of Object-Oriented Queries, Proc SIGMOD International Conference on Management of Data, San Diego, California, Jun. 2-5, 1992, SIGMOD Record, vol. 21, Issue 2, Jun., 1992; Stonebraker, M., Editor;. Other resources are available, e.g., from Microsoft, IBM, Sun and other software development companies.

[0291] Using the tools described above, users of the reagents, methods and database as discovery or diagnostic tools can query a centrally located database with expression and subject data. Each submission of data adds to the sum of expression and subject information in the database. As data is added, a new correlation statistical analysis is automatically run that incorporates the added clinical and expression data. Accordingly, the predictive accuracy and the types of correlations of the recorded molecular signatures increases as the database grows.

[0292] For example, subjects, such as patients, can access the results of the expression analysis of their leukocyte samples and any accrued knowledge regarding the likelihood of the patient's belonging to any specified diagnostic (or prognostic, or monitoring, or risk group), i.e., their expression profiles, and/or molecular signatures. Optionally, subjects can add to the predictive accuracy of the database by providing additional information to the database regarding diagnoses, test results, clinical or other related events that have occurred since the time of the expression profiling. Such information can be provided to the database via any form of communication, including, but not limited to, the internet. Such data can be used to continually define (and redefine) diagnostic groups. For example, if 1000 patients submit data regarding the occurrence of myocardial infarction over the 5 years since their expression profiling, and 300 of these patients report that they have experienced a myocardial infarction and 700 report that they have not, then the 300 patients define a new "group A." As the algorithm is used to continually query and revise the database, a new diagnostic nucleotide set that differentiates groups A and B (i.e., with and without myocardial infarction within a five year period) is identified. This newly defined nucleotide set is then be used (in the manner described above) as a test that predicts the occurrence of myocardial infarction over a five-year period. While submission directly by the patient is exemplified above, any individual with access and authority to submit the relevant data e.g., the patient's physician, a laboratory technician, a health care or study administrator, or the like, can do so.

[0293] As will be apparent from the above examples, transmission of information via the internet (or via an intranet) is optionally bi-directional. That is, for example, data regarding expression profiles, subject data, and the like are transmitted via a communication system to the database, while information regarding molecular signatures, predictive analysis, and the like, are transmitted from the database to the user. For example, using appropriate configurations of an integrated system including a microarray comprising a diagnostic nucleotide set, a detector linked to a computational device can directly transmit (locally or from a remote workstation at great distance, e.g., hundreds or thousands of miles distant from the database) expression profiles and a corresponding individual identifier to a central database for analysis according to the methods of the disclosure. According to, e.g., the algorithms described above, the individual identifier is assigned to one or more diagnostic (or prognostic, or monitoring, etc.) categories. The results of this classification are then relayed back, via, e.g., the same mode of communication, to a recipient at the same or different internet (or intranet) address.

40 Kits

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[0294] The object disclosed is optionally provided to a user as a kit. Typically, a kit contains one or more diagnostic nucleotide sets of the disclosure. Alternatively, the kit contains the candidate nucleotide library of the disclosure. Most often, the kit contains a diagnostic nucleotide probe set, or other subset of a candidate library, e.g., as a cDNA or antibody microarray packaged in a suitable container. The kit may further comprise, one or more additional reagents, e.g., substrates, labels, primers, for labeling expression products, tubes and/or other accessories, reagents for collecting blood samples, buffers, e.g., erythrocyte lysis buffer, leukocyte lysis buffer, hybridization chambers, cover slips, etc., as well as a software package, e.g., including the statistical methods of the disclosure, e.g., as described above, and a password and/or account number for accessing the compiled database. The kit optionally further comprises an instruction set or user manual detailing preferred methods of using the diagnostic nucleotide sets in the methods of the disclosure. In one embodiment, the kit may include contents useful for the discovery of diagnostic nucleotide sets using microarrays. The kit may include sterile, endotoxin and RNAse free blood collection tubes. The kit may also include alcohol swabs, tourniquet, blood collection set, and/or PBS (phosphate buffer saline; needed when method of example 2 is used to derived mononuclear RNA). The kit may also include cell lysis buffer. The kit may include RNA isolation kit, substrates for labeling of RNA (may vary for various expression profiling techniques). The kit may also include materials for fluorescence microarray expression profiling, including one or more of the following: reverse transcriptase and 10x RT buffer, T7(dT)24 primer (primer with T7 promoter at 5' end), DTT, deoxynucleotides, optionally 100mM each, RNAse inhibitor, second strand cDNA buffer, DNA polymerase, Rnase H, T7 RNA polymerase ribonucleotides, in vitro transcription buffer,

and/or Cy3 and Cy5 labeled ribonucleotides. The kit may also include microarrays containing candidate gene libraries, cover slips for slides, and/or hybridization chambers. The kit may further include software package for identification of diagnostic gene set from data, that contains statistical methods, and/or allows alteration in desired sensitivity and specificity of gene set. The software may further facilitate access to and data analysis by centrally a located database server. The software may further include a password and account number to access central database server. In addition, the kit may include a kit user manual.

[0295] In another embodiment, the kit may include contents useful for the application of diagnostic nucleotide sets using microarrays. The kit may include sterile, endotoxin and/or RNAse free blood collection tubes. The kit may also include, alcohol swabs, tourniquet, and/or a blood collection set. The kit may further include PBS (phosphate buffer saline; needed when method of example 2 is used to derived mononuclear RNA), cell lysis buffer, and/or an RNA isolation kit. In addition, the kit may include substrates for labeling of RNA (may vary for various expression profiling techniques). For fluorescence microarray expression profiling, components may include reverse transcriptase and 10x RT buffer, T7 (dT)24 primer (primer with T7 promoter at 5' end), DTT, deoxynucleotides (optionally 100mM each), RNAse inhibitor, second strand cDNA buffer, DNA polymerase, Rnase H, T7 RNA polymerase, ribonucleotides, in vitro transcription buffer, and/or Cy3 and Cy5 labeled ribonucleotides. The kit may further include microarrays containing candidate gene libraries. The kit may also include cover slips for slides, and/or hybridization chambers. The kit may include a software package for identification of diagnostic gene set from data. The software package may contain statistical methods, allow alteration in desired sensitivity and specificity of gene set, and/or facilitate access to and data analysis by centrally located database server. The software package may include a password and account number to access central database server. In addition, the kit may include a kit user manual.

[0296] In another embodiment, the kit may include contents useful for the application of diagnostic nucleotide sets using real-time PCR. This kit may include terile, endotoxin and/or RNAse free blood collection tubes. The kit may further include alcohol swabs, tourniquet, and/or a blood collection set. The kit may also include PBS (phosphate buffer saline; needed when method of example 2 is used to derived mononuclear RNA). In addition, the kit may include cell lysis buffer and/or an RNA isolation kit. The kit may laso include substrates for real time RT-PCR, which may vary for various realtime PCR techniques, including poly dT primers, random hexamer primers, reverse Transcriptase and RT buffer, DTT, deoxynucleotides 100 mM, RNase H, primer pairs for diagnostic and control gene set, 10x PCR reaction buffer, and/or Tag DNA polymerase. The kit may also include fluorescent probes for diagnostic and control gene set (alternatively, fluorescent dye that binds to only double stranded DNA). The kit may further include reaction tubes with or without barcode for sample tracking, 96-well plates with barcode for sample identification, one barcode for entire set, or individual barcode per reaction tube in plate. The kit may also include a software package for identification of diagnostic gene set from data, and /or statistical methods. The software package may allow alteration in desired sensitivity and specificity of gene set, and/or facilitate access to and data analysis by centrally located database server. The kit may include a password and account number to access central database server. Finally, the kit may include a kit user manual.

[0297] This invention will be better understood by reference to the following non-limiting Examples:

LIST OF EXAMPLE TITLES

[0298]

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- Example 1: Preparation of a leukocyte cDNA array comprising a candidate gene library
- Example 2: Preparation of RNA from mononuclear cells for expression profiling
- Example 3: Preparation of Universal Control RNA for use in leukocyte expression profiling
- Example 4. RNA Labeling and hybridization to a leukocyte cDNA array of candidate nucleotide sequences.
- 45 Example 5: Clinical study for the Identification of diagnostic gene sets useful in diagnosis and treatment of Cardiac allograft rejection
 - Example 6: Identification of diagnostic nucleotide sets for kidney and liver allograft rejection
 - Example 7: Identification of diagnostic nucleotide sets for diagnosis of cytomegalovirus
 - Example 8: Design of oligonucleotide probes
- 50 Example 9: Production of an array of 8,000 spotted 50 mer oligonucleotides.
 - Example 10: Identification of diagnostic nucleotide sets for diagnosis of Cardiac Allograft Rejection using microarrays
 - Example 11: Amplification, labeling, and hybridization of total RNA to an oligonucleotide microarray
 - Example 12: Real-time PCR validation of array expression results
 - Example 13: Real-time PCR expression markers of acute allograft rejection
- 55 Example 14: Identification of diagnostic nucleotide sets for diagnosis of Cardiac Allograft Rejection using microarrays
 - Example 15: Correlation and Classification Analysis
 - Example 16: Acute allograft rejection: biopsy tissue gene expression profiling
 - Example 17: Microarray and PCR gene expression panels for diagnosis and monitoring of acute allograft rejection

- Example 18: Assay sample preparation
- Example 19: Allograft rejection diagnostic gene sequence analysis
- Example 20: Detection of proteins expressed by diagnostic gene sequences
- Example 21: Detecting changes in the rate of hematopoiesis

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Examples

Example 1: Preparation of a leukocyte cDNA array comprising a candidate gene

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[0299] Candidate genes and gene sequences for leukocyte expression profiling are identified through methods described elsewhere in this document. Candidate genes are used to obtain or design probes for peripheral leukocyte expression profiling in a variety of ways.

[0300] A cDNA microarray carrying 384 probes was constructed using sequences selected from the initial candidate library. cDNAs is selected from T-cell libraries, PBMC libraries and buffy coat libraries.

96-Well PCR

[0301] Plasmids are isolated in 96-well format and PCR was performed in 96-well format. A master mix is made that contain the reaction buffer, dNTPs, forward and reverse primer and DNA polymerase was made. 99 ul of the master mix was aliquoted into 96-well plate. 1 ul of plasmid (1-2 ng/ul) of plasmid was added to the plate. The final reaction concentration was 10 mM Tris pH 8.3, 3.5 mM MgCl2, 25 mM KCl, 0.4 mM dNTPs, 0.4 uM M13 forward primer, 0.4 M13 reverse primer, and 10 U ofTaq Gold (Applied Biosystems). The PCR conditions were:

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Step 1 95C for 10 min

Step 2 95C for 15 sec

Step 3 56C for 30 sec

Step 4 72C for 2 min 15 seconds

Step 5 go to Step 2 39 times

Step 6 72C for 10 minutes

Step 7 4C for ever.

PCR Purification

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[0302] PCR purification is done in a 96-well format. The Arraylt (Telechem International, Inc.) PCR purification kit is used and the provided protocol was followed without modification. Before the sample is evaporated to dryness, the concentration of PCR products was determined using a spectrophotometer. After evaporation, the samples are resuspended in 1x Micro Spotting Solution (Arraylt) so that the majority of the samples were between 0.2-1.0 ug/ul.

Array Fabrication

[0303] Spotted cDNA microarrays are then made from these PCR products by Arraylt using their protocols, which may be found at the Arraylt website. Each fragment was spotted 3 times onto each array. Candidate genes and gene sequences for leukocyte expression profiling are identified through methods described elsewhere in this document. Those candidate genes are used for peripheral leukocyte expression profiling. The candidate libraries can used to obtain or design probes for expression profiling in a variety of ways.

[0304] Oligonucleotide probes are prepared using the gene sequences of Table 2, and the sequence listing. Oligo probes are designed on a contract basis by various companies (for example, Compugen, Mergen, Affymetrix, Telechem), or designed from the candidate sequences using a variety of parameters and algorithms as indicated at located at the MIT web site. Briefly, the length of the oligonucleotide to be synthesized is determined, preferably greater than 18 nucleotides, generally 18-24 nucleotides, 24-70 nucleotides and, in some circumstances, more than 70 nucleotides. The sequence analysis algorithms and tools described above are applied to the sequence to mask repetitive elements, vector sequences and low complexity sequences. Oligonucleotides are selected that are specific to the candidate nucleotide sequence (based on a Blast n search of the oligonucleotide sequence in question against gene sequences databases, such as the Human Genome Sequence, UniGene, dbEST or the non-redundant database at NCBI), and have <50% G content and 25-70% G+C content. Desired oligonucleotides are synthesized using well-known methods and apparatus, or ordered from a company (for example Sigma). Oligonucleotides are spotted onto microarrays. Alternatively, oligonu-

cleotides are synthesized directly on the array surface, using a variety of techniques (Hughes et al. 2001, Yershov et al. 1996, Lockhart et al 1996).

Example 2: Preparation of RNA from mononuclear cells for expression profiling

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[0305] Blood was isolated from the subject for leukocyte expression profiting using the following methods: Two tubes were drawn per patient. Blood was drawn from either a standard peripheral venous blood draw or directly from a large-bore intra-arterial or intravenous catheter inserted in the femoral artery, femoral vein, subclavian vein or internal jugular vein. Care was taken to avoid sample contamination with heparin from the intravascular catheters, as heparin can interfere with subsequent RNA reactions. For each tube, 8 ml of whole blood was drawn into a tube (CPT, Becton-Dickinson order #362753) containing the anticoagulant Citrate, 25°C density gradient solution (e.g. Ficoll, Percoll) and a polyester gel barrier that upon centrifugation was permeable to RBCs and granulocytes but not to mononuclear cells. The tube was inverted several times to mix the blood with the anticoagulant. The tubes were centrifuged at 1750x in a swing-out rotor at room temperature for 20 minutes. The tubes were removed from the centrifuge and inverted 5-10 times to mix the plasma with the mononuclear cells, while trapping the RBCs and the granulocytes beneath the gel barrier. The plasma/mononuclear cell mix was decanted into a 15ml tube and 5ml of phosphate-buffered saline (PBS) is added. The 15ml tubes were spun for 5 minutes at 1750xg to pellet the cells. The supernatant was discarded and 1.8 ml of RLT lysis buffer is added to the mononuclear cell pellet. The buffer and cells were pipetted up and down to ensure complete lysis of the pellet. The cell lysate was frozen and stored until it is convenient to proceed with isolation of total RNA.

[0306] Total RNA was purified from the lysed mononuclear cells using the Qiagen Rneasy Miniprep kit, as directed by the manufacturer (10/99 version) for total RNA isolation, including homogenization (Qiashredder columns) and on-column DNase treatment. The purified RNA was eluted in 50ul of water. The further use of RNA prepared by this method is described in Examples 10 and 11. Some samples were prepared by a different protocol, as follows:

Two 8 ml blood samples were drawn from a peripheral vein into a tube (CPT, Becton-Dickinson order #362753) containing anticoagulant (Citrate), 25°C density gradient solution (Ficoll) and a polyester gel barrier that upon centrifugation is permeable to RBCs and granulocytes but not to mononuclear cells. The mononuclear cells and plasma remained above the barrier while the RBCs and granulocytes were trapped below. The tube was inverted several times to mix the blood with the anticoagulant, and the tubes were subjected to centrifugation at 1750xg in a swing-out rotor at room temperature for 20 min. The tubes were removed from the centrifuge, and the clear plasma layer above the cloudy mononuclear cell layer was aspirated and discarded. The cloudy mononuclear cell layer was aspirated, with care taken to rinse all of the mononuclear cells from the surface of the gel barrier with PBS (phosphate buffered saline). Approximately 2 mls of mononuclear cell suspension was transferred to a 2ml microcentrifuge tube, and centrifuged for 3min. at 16,000 rpm in a microcentrifuge to pellet the cells. The supernatant was discarded and 1.8 ml of RLT lysis buffer (Qiagen) were added to the mononuclear cell pellet, which lysed the cells and inactivated Rnases. The cells and lysis buffer were pipetted up and down to ensure complete lysis of the pellet. Cell lysate was frozen and stored until it was convenient to proceed with isolation of total RNA.

[0307] RNA samples were isolated from 8 mL of whole blood. Yields ranged from 2 ug to 20ug total RNA for 8mL blood. A260/A280 spectrophotometric ratios were between 1.6 and 2.0, indicating purity of sample. 2ul of each sample were run on an agarose gel in the presence of ethidium bromide. No degradation of the RNA sample and no DNA contamination was visible.

[0308] In some cases, specific subsets of mononuclear cells were isolated from peripheral blood of human subjects. When this was done, the StemSep cell separation kits (manual version 6.0.0) were used from StemCell Technologies (Vancouver, Canada). This same protocol can be applied to the isolation of T cells, CD4 T cells, CD8 T cells, B cells, monocytes, NK cells and other cells. Isolation of cell types using negative selection with antibodies may be desirable to avoid activation of target cells by antibodies.

Example 3: Preparation of Universal Control RNA for use in leukocyte expression profiling

[0309] Control RNA was prepared using total RNA from Buffy coats and/or total RNA from enriched mononuclear cells isolated from Buffy coats, both with and without stimulation with ionomycin and PMA. The following control RNAs were prepared:

55 Control 1: Buffy Coat Total RNA

Control 2: Mononuclear cell Total RNA

Control 3: Stimulated buffy coat Total RNA

Control 4: Stimulated mononuclear Total RNA

Control 5: 50% Buffy coat Total RNA / 50% Stimulated buffy coat Total RNA

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Control 6: 50% Mononuclear cell Total RNA / 50% Stimulated Mononuclear Total RNA

[0310] Some samples were prepared using the following protocol: Buffy coats from 38 individuals were obtained from Stanford Blood Center. Each buffy coat is derived from -350 mL whole blood from one individual. 10 ml buffy coat was removed from the bag, and placed into a 50 ml tube. 40 ml of Buffer EL (Qiagen) was added, the tube was mixed and placed on ice for 15 minutes, then cells were pelleted by centrifugation at 2000xg for 10 minutes at 4°C. The supernatant was decanted and the cell pellet was re-suspended in 10 ml of Qiagen Buffer EL. The tube was then centrifuged at 2000xg for 10 minutes at 4°C. The cell pellet was then re-suspended in 20 ml TRIZOL (GibcoBRL) per Buffy coat sample, the mixture was shredded using a rotary homogenizer, and the lysate was then frozen at -80°C prior to proceeding to RNA isolation.

[0311] Other control RNAs were prepared from enriched mononuclear cells prepared from Buffy coats. Buffy coats from Stanford Blood Center were obtained, as described above. 10 ml buffy coat was added to a 50 ml polypropylene tube, and 10 ml of phosphate buffer saline (PBS) was added to each tube. A polysucrose (5.7 g/dL) and sodium diatrizoate (9.0 g/dL) solution at a 1.077 +/-0.0001 g/ml density solution of equal volume to diluted sample was prepared (Histopaque 1077, Sigma cat. no 1077-1). This and all subsequent steps were performed at room temperature. 15 ml of diluted buffy coat/PBS was layered on top of 15 ml of the histopaque solution in a 50 ml tube. The tube was centrifuged at 400xg for 30 minutes at room temperature. After centrifugation, the upper layer of the solution to within 0.5 cm of the opaque interface containing the mononuclear cells was discarded. The opaque interface was transferred into a clean centrifuge tube. An equal volume of PBS was added to each tube and centrifuged at 350xg for 10 minutes at room temperature. The supernatant was discarded. 5 ml of Buffer EL (Qiagen) was used to resuspend the remaining cell pellet and the tube was centrifuged at 2000xg for 10 minutes at room temperature. The supernatant was discarded. The pellet was resuspended in 20 ml of TRIZOL (GibcoBRL) for each individual buffy coat that was processed. The sample was homogenized using a rotary homogenizer and frozen at -80C until RNA was isolated. RNA was isolated from frozen lysed Buffy coat samples as follows: frozen samples were thawed, and 4 ml of chloroform was added to each buffy coat sample. The sample was mixed by vortexing and centrifuged at 2000xg for 5 minutes. The aqueous layer was moved to new tube and then repurified by using the RNeasy Maxi RNA clean up kit, according to the manufacturer's instruction (Qiagen, PN 75162). The yield, purity and integrity were assessed by spectrophotometer and gel electrophoresis. Some samples were prepared by a different protocol, as follows. The further use of RNA prepared using this protocol is described in Example 11.

[0312] 50 whole blood samples were randomly selected from consented blood donors at the Stanford Medical School Blood Center. Each buffy coat sample was produced from ~350 mL of an individual's donated blood. The whole blood sample was centrifuged at ~4,400 x g for 8 minutes at room temperature, resulting in three distinct layers: a top layer of plasma, a second layer of buffy coat, and a third layer of red blood cells. 25 ml of the buffy coat fraction was obtained and diluted with an equal volume of PBS (phosphate buffered saline). 30 ml of diluted buffy coat was layered onto 15 ml of sodium diatrizoate solution adjusted to a density of 1.077+/-0.001 g/ml (Histopaque 1077, Sigma) in a 50mL plastic tube. The tube was spun at 800 g for 10 minutes at room temperature. The plasma layer was removed to the 30 ml mark on the tube, and the mononuclear cell layer removed into a new tube and washed with an equal volume of PBS, and collected by centrifugation at 2000 g for 10 minutes at room temperature. The cell pellet was resuspended in 10 ml of Buffer EL (Qiagen) by vortexing and incubated on ice for 10 minutes to remove any remaining erthythrocytes. The mononuclear cells were spun at 2000 g for 10 minutes at 4 degrees Celsius. The cell pellet was lysed in 25 ml of a phenol/guanidinium thiocyanate solution (TRIZOL Reagent, Invitrogen). The sample was homogenized using a Power-Gene 5 rotary homogenizer (Fisher Scientific) and Omini disposable generator probes (Fisher Scientific). The Trizol lysate was frozen at -80 degrees C until the next step.

[0313] The samples were thawed out and incubated at room temperature for 5 minutes. 5 ml chloroform was added to each sample, mixed by vortexing, and incubated at room temperature for 3 minutes. The aqueous layers were transferred to new 50 ml tubes. The aqueous layer containing total RNA was further purified using the Qiagen RNeasy Maxi kit (PN 75162), per the manufacturer's protocol (October 1999). The columns were eluted twice with 1 ml Rnase-free water, with a minute incubation before each spin. Quantity and quality of RNA was assessed using standard methods.
 Generally, RNA was isolated from batches of 10 buffy coats at a time, with an average yield per buffy coat of 870 μg, and an estimated total yield of 43.5 mg total RNA with a 260/280 ratio of 1.56 and a 28S/18S ratio of 1.78.

[0314] Quality of the RNA was tested using the Agilent 2100 Bioanalyzer using RNA 6000 microfluidics chips. Analysis of the electrophorgrams from the Bioanalyzer for five different batches demonstrated the reproducibility in quality between the batches.

[0315] Total RNA from all five batches were combined and mixed in a 50 ml tube, then aliquoted as follows: 2 x 10 ml aliquots in 15 ml tubes, and the rest in 100 μl aliquots in 1.5 ml microcentrifuge tubes. The aliquots gave highly reproducible results with respect to RNA purity, size and integrity. The RNA was stored at -80°C.

Test hybridization of Reference RNA.

[0316] When compared with BC38 and Stimulated mononuclear reference samples, the R50 performed as well, if not better than the other reference samples as shown in Figure 3. In an analysis of hybridizations, where the R50 targets were fluorescently labeled with Cy-5 using methods described herein and the amplified and labeled aRNA was hybridized (as in example 11) to the olignoucleotide array described in example 9. The R50 detected 97.3% of probes with a Signal to Noise ratio (S/N) of greater than three and 99.9 % of probes with S/N greater than one.

Example 4. RNA Labeling and hybridization to a leukocyte cDNA array of candidate nucleotide sequences.

Comparison of Guanine-Silica to Acid-Phenol RNA Purification (GSvsAP)

[0317] These data are from a set of 12 hybridizations designed to identify differences between the signal strength from two different RNA purification methods. The two RNA methods used were guanidine-silica (GS, Qiagen) and acid-phenol (AP, Trizol, Gibco BRL). Ten tubes of blood were drawn from each of four people. Two were used for the AP prep, the other eight were used for the GS prep. The protocols for the leukocyte RNA preps using the AP and GS techniques were completed as described here:

Guanidine-silica (GS) method:

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[0318] For each tube, 8ml blood was drawn into a tube containing the anticoagulant Citrate, 25°C density gradient solution and a polyester gel barrier that upon centrifugation is permeable to RBCs and granulocytes but not to mononuclear cells. The mononuclear cells and plasma remained above the barrier while the RBCs and granulocytes were trapped below. CPT tubes from Becton-Dickinson (#362753) were used for this purpose. The tube was inverted several times to mix the blood with the anticoagulant. The tubes were immediately centrifuged @1750xg in a swinging bucket rotor at room temperature for 20 min. The tubes were removed from the centrifuge and inverted 5-10 times. This mixed the plasma with the mononuclear cells, while the RBCs and the granulocytes remained trapped beneath the gel barrier. The plasma/mononuclear cell mix was decanted into a 15ml tube and 5ml of phosphate-buffered saline (PBS) was added. The 15ml tubes are spun for 5 minutes at 1750xg to pellet the cells. The supernatant was discarded and 1.8 ml of RLT lysis buffer (guanidine isothyocyanate) was added to the mononuclear cell pellet. The buffer and cells were pipetted up and down to ensure complete lysis of the pellet. The cell lysate was then processed exactly as described in the Qiagen Rneasy Miniprep kit protocol (10/99 version) for total RNA isolation (including steps for homogenization (Qiashredder columns) and on-column DNase treatment. The purified RNA was eluted in 50ul of water.

35 <u>Acid-phenol (AP) method:</u>

[0319] For each tube, 8ml blood was drawn into a tube containing the anticoagulant Citrate, 25°C density gradient solution and a polyester gel barrier that upon centrifugation is permeable to RBCs and granulocytes but not to mononuclear cells. The mononuclear cells and plasma remained above the barrier while the RBCs and granulocytes were trapped below. CPT tubes from Becton-Dickinson (#362753) were used for this purpose. The tube was inverted several times to mix the blood with the anticoagulant. The tubes were immediately centrifuged @1750xg in a swinging bucket rotor at room temperature for 20 min. The tubes were removed from the centrifuge and inverted 5-10 times. This mixed the plasma with the mononuclear cells, while the RBCs and the granulocytes remained trapped beneath the gel barrier. The plasma/mononuclear cell mix was decanted into a 15ml tube and 5ml of phosphate-buffered saline (PBS) was added. The 15ml tubes are spun for 5 minutes @1750xg to pellet the cells. The supernatant was discarded and the cell pellet was lysed using 0.6 mL Phenol/guanidine isothyocyanate (e.g. Trizol reagent, GibcoBRL). Subsequent total RNA isolation proceeded using the manufacturers protocol.

[0320] RNA from each person was labeled with either Cy3 or Cy5, and then hybridized in pairs to the mini-array. For instance, the first array was hybridized with GS RNA from one person (Cy3) and GS RNA from a second person (Cy5). [0321] Techniques for labeling and hybridization for all experiments discussed here were completed as detailed above. Arrays were prepared as described in example 1.

[0322] RNA isolated from subject samples, or control Buffy coat RNA, were labeled for hybridization to a cDNA array. Total RNA (up to 100 μ g) was combined with 2 μ l of 100 μ M solution of an Oligo (dT)12-18 (GibcoBRL) and heated to 70°C for 10 minutes and place on ice. Reaction buffer was added to the tube, to a final concentration of 1xRT buffer (GibcoBRL), 10 mM DTT (GibcoBRL), 0.1 mM unlabeled dATP, dTTP, and dGTP, and 0.025 mM unlabeled dCTP, 200 pg of CAB (A. thaliana photosystem I chlorophyll a/b binding protein), 200 pg of RCA (A. thaliana RUBISCO activase), 0.25 mM of Cy-3 or Cy-5 dCTP, and 400 U Superscript II RT (GibcoBRL).

[0323] The volumes of each component of the labeling reaction were as follows: 20 μl of 5xRT buffer; 10 μl of 100

mM DTT; 1 μ l of 10 mM dNTPs without dCTP; 0.5 μ l of 5 mM CTP; 1 μ l of H20; 0.02 μ l of 10 ng/ μ l CAB and RCA; I μ l of 40 Units/µl RNAseOUT Recombinatnt Ribonuclease Inhibitor (GibcoBRL); 2.5 µl of 1.0 mM Cy-3 or Cy-5 dCTP; and 2.0 μl of 200 Units/μl of Superscript II RT. The sample was vortexed and centrifuged. The sample was incubated at 4°C for 1 hour for first strand cDNA synthesis, then heated at 70°C for 10 minutes to quench enzymatic activity. 1 μ l of 10 mg/ml of Rnase A was added to degrade the RNA strand, and the sample was incubated at 37°C for 30 minutes. Next, the Cy-3 and Cy-5 cDNA samples were combined into one tube. Unincorporated nucleotides were removed using QIAquick RCR purification protocol (Qiagen), as directed by the manufacturer. The sample was evaporated to dryness and resuspended in 5 μ l of water. The sample was mixed with hybridization buffer containing 5xSSC, 0.2% SDS, 2 mg/ml Cot-1 DNA (GibcoBRL), 1 mg/ml yeast tRNA (GibcoBRL), and 1.6 ng/ μ l poly dA40-60 (Pharmacia). This mixture was placed on the microarray surface and a glass cover slip was placed on the array (Coming). The microarray glass slide was placed into a hybridization chamber (Arrraylt). The chamber was then submerged in a water bath overnight at 62° C. The microarray was removed from the cassette and the cover slip was removed by repeatedly submerging it to a wash buffer containing 1xSSC, and 0.1 % SDS. The microarray slide was washed in 1xSSC/0.1% SDS for 5 minutes. The slide was then washed in 0.1%SSC/0.1% SDS for 5 minutes. The slide was finally washed in 0.1xSSC for 2 minutes. The slide was spun at 1000 rpm for 2 minutes to dry out the slide, then scanned on a microarray scanner (Axon Instruments, Union City, CA.).

[0324] Six hybridizations with 20 μ g of RNA were performed for each type of RNA preparation (GS or AP). Since both the Cy3 and the Cy5 labeled RNA are from test preparations, there are six data points for each GS prepped, Cy3-labeled RNA and six for each GS-prepped, Cy5-labeled RNA. The mini array hybridizations were scanned on and Axon Instruments scanner using GenPix 3.0 software. The data presented were derived as follows. First, all features flagged as "not found" by the software were removed from the dataset for individual hybridizations. These features are usually due to high local background or other processing artifacts. Second, the median fluorescence intensity minus the background fluorescence intensity was used to calculate the mean background subtracted signal for each dye for each hybridization. In Figure 3, the mean of these means across all six hybridizations is graphed (n=6 for each column). The error bars are the SEM. This experiment shows that the average signal from AP prepared RNA is 47% of the average signal from GS prepared RNA for both Cy3 and Cy5.

Generation of expression data for leukocyte genes from peripheral leukocyte samples

30 **[0325]** Six hybridizations were performed with RNA purified from human blood leukocytes using the protocols given above. Four of the six were prepared using the GS method and 2 were prepared using the AP method. Each preparation of leukocyte RNA was labeled with Cy3 and 10 μg hybridized to the mini-array. A control RNA was batch labeled with Cy5 and 10 μg hybridized to each mini-array together with the Cy3-labeled experimental RNA.

[0326] The control RNA used for these experiments was Control 1: Buffy Coat RNA, as described above. The protocol for the preparation of that RNA is reproduced here:

Buffy Coat RNA Isolation:

[0327] Buffy coats were obtained from Stanford Blood Center (in total 38 individual buffy coats were used. Each buffy coat is derived from -350 mL whole blood from one individual. 10 ml buffy coat was taken and placed into a 50 ml tube and 40 ml of a hypoclorous acid (HOCI) solution (Buffer EL from Qiagen) was added. The tube was mixed and placed on ice for 15 minutes. The tube was then centrifuged at 2000xg for 10 minutes at 4°C. The supernatant was decanted and the cell pellet was re-suspended in 10 ml of hypochlorous acid solution (Qiagen Buffer EL). The tube was then centrifuged at 2000xg for 10 minutes at 4°C. The cell pellet was then re-suspended in 20 ml phenol/guanidine thiocyanate solution (TRIZOL from GibcoBRL) for each individual buffy coat that was processed. The mixture was then shredded using a rotary homogenizer. The lysate was then frozen at -80°C prior to proceeding to RNA isolation.

[0328] The arrays were then scanned and analyzed on an Axon Instruments scanner using GenePix 3.0 software. The data presented were derived as follows. First, all features flagged as "not found" by the software were removed from the dataset for individual hybridizations. Second, control features were used to normalize the data for labeling and hybridization variability within the experiment. The control features are cDNA for genes from the plant, Arabidopsis thaliana, that were included when spotting the mini-array. Equal amounts of RNA complementary to two of these cDNAs were added to each of the samples before they were labeled. A third was pre-labeled and equal amounts were added to each hybridization solution before hybridization. Using the signal from these genes, we derived a normalization constant (L_i) according to the following formula:

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$$L_{j} = \frac{\sum_{i=1}^{N} BGSS_{j,i}}{\sum_{i=1}^{N} BGSS_{j,i}}$$

$$\frac{\sum_{j=1}^{K} \sum_{i=1}^{N} BGSS_{j,i}}{N}$$

$$K$$

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where BGSS_i is the signal for a specific feature as identified in the GenePix software as the median background subtracted signal for that feature, N is the number of A. thaliana control features, K is the number of hybridizations, and L is the normalization constant for each individual hybridization. Using the formula above, the mean over all control features of a particular hybridization and dye (eg Cy3) was calculated. Then these control feature means for all Cy3 hybridizations were averaged. The control feature mean in one hybridization divided by the average of all hybridizations gives a normalization constant for that particular Cy3 hybridization.

[0329] The same normalization steps were performed for Cy3 and Cy5 values, both fluorescence and background. Once normalized, the background Cy3 fluorescence was subtracted from the Cy3 fluorescence for each feature. Values less than 100 were eliminated from further calculations since low values caused spurious results.

[0330] Figure 4 shows the average background subtracted signal for each of nine leukocyte-specific genes on the mini array. This average is for 3-6 of the above-described hybridizations for each gene. The error bars are the SEM.

[0331] The ratio of Cy3 to Cy5 signal is shown for a number of genes. This ratio corrects for variability among hybridizations and allows comparison between experiments done at different times. The ratio is calculated as the Cy3 background subtracted signal divided by the Cy5 background subtracted signal. Each bar is the average for 3-6 hybridizations. The error bars are SEM.

[0332] Together, these results show that we can measure expression levels for genes that are expressed specifically in sub-populations of leukocytes. These expression measurements were made with only 10 μ g of leukocyte total RNA that was labeled directly by reverse transcription. The signal strength can be increased by improved labeling techniques that amplify either the starting RNA or the signal fluorescence. In addition, scanning techniques with higher sensitivity can be used.

Genes in Figures 4 and 5:

Concestinatinguites 4 and 6.					
Gene Name/Description	GenBank Accession Number	Gene Name Abbreviation			
T cell-specific tyrosine kinase Mrna	L10717	TKTCS			
Interleukin I alpha (IL 1) mRNA, complete cds	NM_000575	IL1A			
T-cell surface antigen CD2 (T11) mRNA, complete cds	M14362	CD2			
Interleukin-13 (IL-13) precursor gene, complete cds	U31120	IL-13			
Thymocyte antigen CD1a mRNA, complete cds	M28825	CD1a			
CD6 mRNA for T cell glycoprotein CDS	NM_006725	CD6			
MHC class II HLA-DQA1 mRNA, complete cds	U77589	HLA-DQA1			
Granulocyte colony-stimulating factor	M28170	CD19			
Homo sapiens CD69 antigen	NM_001781	CD69			

Example 5: Clinical study to identify diagnostic gene sets useful in diagnosis and treatment of cardiac allograft recipients

[0333] An observational study was conducted in which a prospective cohort of cardiac transplant recipients were analyzed for associations between clinical events or rejection grades and expression of a leukocyte candidate nucleotide sequence library. Patients were identified at 4 cardiac transplantation centers while on the transplant waiting list or during their routing post-transplant care. All adult cardiac transplant recipients (new or re-transplants) who received an organ at the study center during the study period or within 3 months of the start of the study period were eligible. The first year after transplantation is the time when most acute rejection occurs and it is thus important to study patients during this

period. Patients provided informed consent prior to study procedures.

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[0334] Peripheral blood leukocyte samples were obtained from all patients at the following time points: prior to transplant surgery (when able), the same day as routinely scheduled screening biopsies, upon evaluation for suspected acute rejection (urgent biopsies), on hospitalization for an acute complication of transplantation or immunosuppression, and when Cytomegalovirus (CMV) infection was suspected or confirmed. Samples were obtained through a standard peripheral vein blood draw or through a catheter placed for patient care (for example, a central venous catheter placed for endocardial biopsy). When blood was drawn from a intravenous line, care was taken to avoid obtaining heparin with the sample as it can interfere with downstream reactions involving the RNA. Mononuclear cells were prepared from whole blood samples as described in Example 2. Samples were processed within 2 hours of the blood draw and DNA and serum were saved in addition to RNA. Samples were stored at -80° C or on dry ice and sent to the site of RNA preparation in a sealed container with ample dry ice. RNA was isolated from subject samples as described in Example 2 and hybridized to a candidate library of differentially expressed leukocyte nucleotide sequences, as further described in Example 9-10. Methods used for amplification, labeling, hybridization and scanning are described in Example 11. Analysis of human transplant patient mononuclear cell RNA hybridized to a microarray and identification of diagnostic gene sets is shown in Example 10.

[0335] From each patient, clinical information was obtained at the following time points: prior to transplant surgery (when available), the same day as routinely scheduled screening biopsies, upon evaluation for suspected acute rejection (e.g., urgent biopsies), on hospitalization for an acute complication of transplantation or immunosuppression, and when Cytomegalovirus (CMV) infection was suspected or confirmed. Data was collected directly from the patient, from the patient's medical record, from diagnostic test reports or from computerized hospital databases. It was important to collect all information pertaining to the study clinical correlates (diagnoses and patient events and states to which expression data is correlated) and confounding variables (diagnoses and patient events and states that may result in altered leukocyte gene expression. Examples of clinical data collected are: patient sex, date of birth, date of transplant, race, requirement for prospective cross match, occurrence of pre-transplant diagnoses and complications, indication for transplantation, severity and type of heart disease, history of left ventricular assist devices, all known medical diagnoses, blood type, HLA type, viral serologies (including CMV, Hepatitis B and C, HIV and others), serum chemistries, white and red blood cell counts and differentials, CMV infections (clinical manifestations and methods of diagnosis), occurrence of new cancer, hemodynamic parameters measured by catheterization of the right or left heart (measures of graft function), results of echocardiography, results of coronary angiograms, results of intravascular ultrasound studies (diagnosis of transplant vasculopathy), medications, changes in medications, treatments for rejection, and medication levels. Information was also collected regarding the organ donor, including demographics, blood type, HLA type, results of screening cultures, results of viral serologies, primary cause of brain death, the need for inotropic support, and the organ cold ischemia time.

[0336] Of great importance was the collection of the results of endocardial biopsy for each of the patients at each visit. Biopsy results were all interpreted and recorded using the international society for heart and lung transplantation (ISHLT) criteria, described below. Biopsy pathological grades were determined by experienced pathologists at each center.

ISHLT Criteria

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Grade	Finding	Rejection Severity		
0	No lymphocytic infiltrates	None		
1A	Focal (perivascular or interstitial lymphocytic infiltrates without necrosis)	Borderline mild		
1B	Diffuse but sparse lymphocytic infiltrates without necrosis	Mild		
2	One focus only with aggressive lymphocytic infiltrate and/or myocyte damage	Mild, focal moderate		
3A	Multifocal aggressive lymphocytic infiltrates and/or myocardial damage	Moderate		
3B	Diffuse inflammatory lymphocytic infiltrates with necrosis	Borderline Severe		
4	Diffuse aggressive polymorphous lymphocytic infiltrates with edema hemorrhage and vasculitis, with necrosis	Severe		

[0337] Because variability exists in the assignment of ISHLT grades, it was important to have a centralized and blinded reading of the biopsy slides by a single pathologist. This was arranged for all biopsy slides associated with samples in the analysis. Slides were obtained and assigned an encoded number. A single pathologist then read all slides from all centers and assigned an ISHLT grade. Grades from the single pathologist were then compared to the original grades derived from the pathologists at the study centers. For the purposes of correlation analysis of leukocyte gene expression to biopsy grades, the centralized reading information was used in a variety of ways (see Example 10 for more detail).

In some analyses, only the original reading was used as an outcome. In other analyses, the result from the centralized reader was used as an outcome. In other analyses, the highest of the 2 grades was used. For example, if the original assigned grade was 0 and the centralized reader assigned a 1A, then 1A was the grade used as an outcome. In some analyses, the highest grade was used and then samples associated with a Grade 1A reading were excluded from the analysis. In some analyses, only grades with no disagreement between the 2 readings were used as outcomes for correlation analysis.

[0338] Clinical data was entered and stored in a database. The database was queried to identify all patients and patient visits that meet desired criteria (for example, patients with > grade II biopsy results, no CMV infection and time since transplant < 12 weeks).

[0339] The collected clinical data (disease criteria) is used to define patient or sample groups for correlation of expression data. Patient groups are identified for comparison, for example, a patient group that possesses a useful or interesting clinical distinction, versus a patient group that does not possess the distinction. Examples of useful and interesting patient distinctions that can be made on the basis of collected clinical data are listed here:

- 1. Rejection episode of at least moderate histologic grade, which results in treatment of the patient with additional corticosteroids, anti-T cell antibodies, or total lymphoid irradiation.
 - 2. Rejection with histologic grade 2 or higher.
 - 3. Rejection with histologic grade <2.

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- 4. The absence of histologic rejection and normal or unchanged allograft function (based on hemodynamic measurements from catheterization or on echocardiographic data).
- 5. The presence of severe allograft dysfunction or worsening allograft dysfunction during the study period (based on hemodynamic measurements from catheterization or on echocardiographic data).
- 6. Documented CMV infection by culture, histology, or PCR, and at least one clinical sign or symptom of infection.
- 7. Specific graft biopsy rejection grades
- 8. Rejection of mild to moderate histologic severity prompting augmentation of the patient's chronic immunosuppressive regimen
 - 9. Rejection of mild to moderate severity with allograft dysfunction prompting plasmaphoresis or a diagnosis of "humoral" rejection
 - 10. Infections other than CMV, esp. Epstein Barr virus (EBV)
- 11. Lymphoproliferative disorder (also called, post-transplant lymphoma)
- 12. Transplant vasculopathy diagnosed by increased intimal thickness on intravascular ultrasound (IVUS), angiography, or acute myocardial infarction.
- 13. Graft Failure or Retransplantation
- 14. All cause mortality
- 15. Grade 1A or higher rejection as defined by the initial biopsy reading.
- 16. Grade 1B or higher rejection as defined by the initial biopsy reading.
- 17. Grade 1A or higher rejection as defined by the centralized biopsy reading.
- 18. Grade 1B or higher rejection as defined by the centralized biopsy reading.
- 19. Grade 1A or higher rejection as defined by the highest of the initial and centralized biopsy reading.
- 20. Grade 1B or higher rejection as defined by the highest of the initial and centralized biopsy reading.
- 21. Any rejection > Grade 2 occurring in patient at any time in the post-transplant course.

[0340] Expression profiles of subject samples are examined to discover sets of nucleotide sequences with differential expression between patient groups, for example, by methods describes above and below. Non-limiting examples of patient leukocyte samples to obtain for discovery of various diagnostic nucleotide sets are as follows:

Leukocyte set to avoid biopsy or select for biopsy:

Samples: Grade 0 vs. Grades 1-4

50 Leukocyte set to monitor therapeutic response:

Examine successful vs. unsuccessful drug treatment.

Samples:

Successful: Time 1: rejection, Time 2: drug therapy Time 3: no rejection

Unsuccessful: Time 1: rejection, Time 2: drug therapy; Time 3: rejection

Leukocyte set to predict subsequent acute rejection.

Biopsy may show no rejection, but the patient may develop rejection shortly thereafter. Look at profiles of patients

who subsequently do and do not develop rejection.

Samples:

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Group 1 (Subsequent rejection): Time 1: Grade 0; Time 2: Grade>0 Group 2 (No subsequent rejection): Time 1: Grade 0; Time 2: Grade 0

Focal rejection may be missed by biopsy. When this occurs the patient may have a Grade 0, but actually has rejection. These patients may go on to have damage to the graft etc.

Samples:

Non-rejectors: no rejection over some period of time Rejectors: an episode of rejection over same period

Leukocyte set to diagnose subsequent or current graft failure:

Samples:

Echocardiographic or catheterization data to define worsening function over time and correlate to profiles.

Leukocyte set to diagnose impending active CMV:

Samples:

Look at patients who are CMV IgG positive. Compare patients with subsequent (to a sample) clinical CMV infection verses no subsequent clinical CMV infection.

Leukocyte set to diagnose current active CMV:

Samples:

Analyze patients who are CMV IgG positive. Compare patients with active current clinical CMV infection vs. no active current CMV infection.

[0341] Upon identification of a nucleotide sequence or set of nucleotide sequences that distinguish patient groups with a high degree of accuracy, that nucleotide sequence or set of nucleotide sequences is validated, and implemented as a diagnostic test. The use of the test depends on the patient groups that are used to discover the nucleotide set. For example, if a set of nucleotide sequences is discovered that have collective expression behavior that reliably distinguishes patients with no histological rejection or graft dysfunction from all others, a diagnostic is developed that is used to screen patients for the need for biopsy. Patients identified as having no rejection do not need biopsy, while others are subjected to a biopsy to further define the extent of disease. In another example, a diagnostic nucleotide set that determines continuing graft rejection associated with myocyte necrosis (> grade I) is used to determine that a patient is not receiving adequate treatment under the current treatment regimen. After increased or altered immunosuppressive therapy, diagnostic profiling is conducted to determine whether continuing graft rejection is progressing. In yet another example, a diagnostic nucleotide set(s) that determine a patient's rejection status and diagnose cytomegalovirus infection is used to balance immunosuppressive and anti-viral therapy.

[0342] The methods of this example are also applicable to cardiac xenograft monitoring.

50 Example 6: Identification of diagnostic nucleotide sets for kidney and liver allograft rejection

[0343] Diagnostic tests for rejection are identified using patient leukocyte expression profiles to identify a molecular signature correlated with rejection of a transplanted kidney or liver. Blood, or other leukocyte source, samples are obtained from patients undergoing kidney or liver biopsy following liver or kidney transplantation, respectively. Such results reveal the histological grade, i.e., the state and severity of allograft rejection. Expression profiles are obtained from the samples as described above, and the expression profile is correlated with biopsy results. In the case of kidney rejection, clinical data is collected corresponding to urine output, level of creatine clearance, and level of serum creatine (and other markers of renal function). Clinical data collected for monitoring liver transplant rejection includes, biochemical

characterization of serum markers of liver damage and function such as SGOT, SGPT, Alkaline phosphatase, GGT, Bilirubin, Albumin and Prothrombin time.

[0344] Leukocyte nucleotide sequence expression profiles are collected and correlated with important clinical states and outcomes in renal or hepatic transplantation. Examples of useful clinical correlates are given here:

- 1. Rejection episode of at least moderate histologic grade, which results in treatment of the patient with additional corticosteriods, anti-T cell antibodies, or total lymphoid irradiation.
- 2. The absence of histologic rejection and normal or unchanged allograft function (based on tests of renal or liver function listed above).
- 3. The presence of severe allograft dysfunction or worsening allograft dysfunction during the study period (based on tests of renal and hepatic function listed above).
- 4. Documented CMV infection by culture, histology, or PCR, and at least one clinical sign or symptom of infection.
- 5. Specific graft biopsy rejection grades
- 6. Rejection of mild to moderate histologic severity prompting augmentation of the patient's chronic immunosuppressive regimen
- 7. Infections other than CMV, esp. Epstein Barr virus (EBV)
- 8. Lymphoproliferative disorder (also called, post-transplant lymphoma)
- 9. Graft Failure or Retransplantation
- 10. Need for hemodialysis or other renal replacement therapy for renal transplant patients.
- 11. Hepatic encephalopathy for liver transplant recipients.
- 12. All cause mortality

[0345] Subsets of the candidate library (or of a previously identified diagnostic nucleotide set), are identified, according to the above procedures, that have predictive and/or diagnostic value for kidney or liver allograft rejection.

Example 7: Identification of a diagnostic nucleotide set for diagnosis of cytomegalovirus

[0346] Cytomegalovirus is a very important cause of disease in immunocompromised patients, for example, transplant patients, cancer patients, and AIDS patients. The virus can cause inflammation and disease in almost any tissue (particularly the colon, lung, bone marrow and retina). It is increasingly important to identify patients with current or impending clinical CMV disease, particularly when immunosuppressive drugs are to be used in a patient, e.g. for preventing transplant rejection. Leukocytes are profiled in patients with active CMV, impending CMV, or no CMV. Expression profiles correlating with diagnosis of active or impending CMV are identified. Subsets of the candidate library (or a previously identified diagnostic nucleotide set) are identified, according to the above procedures that have predictive value for the diagnosis of active or impending CMV. Diagnostic nucleotide set(s) identified with predictive value for the diagnosis of active or impending CMV may be combined, or used in conjunction with, cardiac, liver and/or kidney allograft-related diagnostic gene set(s) (described in Examples 6 and 10).

[0347] In addition, or alternatively, CMV nucleotide sequences are obtained, and a diagnostic nucleotide set is designed using CMV nucleotide sequence. The entire sequence of the organism is known and all CMV nucleotide sequences can be isolated and added to the library using the sequence information and the approach described below. Known expressed genes are preferred. Alternatively, nucleotide sequences are selected to represent groups of CMV genes that are coordinately expressed (immediate early genes, early genes, and late genes) (Spector et al. 1990, Stamminger et al. 1990).

[0348] Oligonucleotides were designed for CMV genes using the oligo design procedures of Example 8. Probes were designed using the 14 gene sequences shown here and were included on the array described in example 9:

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		HCMVTRL2 (IRL2)	18932240
5		HCMVTRL7 (IRL7)	complement(65956843)
		HCMVUL21	complement(2649727024)
		HCMVUL27	complement(3283134657)
		HCMVUL33	4325144423
		HCMVUL54	complement(7690380631)
10	Cytomegalovirus (CMV) Accession	HCMVUL75	complement(107901110132)
	#X17403	HCMVUL83	complement(119352121037)
		HCMVUL106	complement(154947155324)
		HCMVUL109	complement(157514157810)
15		HCMVUL113	161503162800
		HCMVUL122	complement(169364170599)
		HCMVUL123 (last exon at 3'-end)	complement(171006172225)
		HCMVUS28	219200220171

[0349] Diagnostic nucleotide set(s) for expression of CMV genes is used in combination with diagnostic leukocyte nucleotide sets for diagnosis of other conditions, e.g. organ allograft rejection.

[0350] Using the techniques described in example 2 mononuclear samples from 180 cardiac transplant recipients (enrolled in the study described in Example 5) were used for expression profiling with the leukocyte arrays. Of these samples 15 were associated with patients who had a diagnosis of primary or reactivation CMV made by culture, PCR or any specific diagnostic test.

[0351] After preparation of RNA, amplification, labeling, hybridization, scanning, feature extraction and data processing were done as described in Example 11 using the oligonucleotide microarrays described in Example 9.

[0352] The resulting log ratio of expression of Cy3 (patient sample)/ Cy5 (R50 reference RNA) was used for analysis. Significance analysis for microarrays (SAM, Tusher 2001, see Example 15) was applied to determine which genes were most significantly differentially expressed between these 15 CMV patients and the 165 non-CMV patients 12 genes were identified with a 0% FDR and 6 with a 0.1% FDR and are listed in Table 2. Some genes are represented by more than one oligonucleotide on the array and for 2 genes, multiple oligonucleotides from the same gene are called significant (SEQ IDs: 3061, 3064: eomesodermin and 3031, 3040,104, 2736: small inducible cytokine A4).

[0353] Clinical variables were also included in the significance analysis. For example, the white blood cell count and the number of weeks post transplant (for the patient at the time the sample was obtained) were available for most of the 180 samples. The log of these variables was taken and the variables were then used in the significance analysis described above with the gene expression data. Both the white blood cell count (0.1% FDR) and the weeks post transplant (0% FDR) appeared to correlate with CMV status. CMV patients were more likely to have samples associated with later post transplant data and the lower white blood cell counts.

[0354] These genes and variables can be used alone or in association with other genes or variables or with other genes to build a diagnostic gene set or a classification algorithm using the approaches described herein.

[0355] Primers for real-time PCR validation were designed for some of these genes as described in Example 13 and listed in Table 2C and the sequence listing. Using the methods described in example 13, primers for Granzyme B were designed and used to validate expression findings from the arrays. 6 samples were tested (3 from patients with CMV and 3 from patients without CMV). The gene was found to be differentially expressed between the patients with and without CMV (see example 13 for full description). This same approach can be used to validate other diagnostic genes by real-time PCR. Diagnostic nucleotide sets can also be identified for a variety of other viral diseases (Table 1) using this same approach.

[0356] cDNA microarrays may be used to monitor viral expression. In addition, these methods may be used to monitor other viruses, such as Epstein-Barr virus, Herpes Simplex 1 and vesicular stomatitis virus.

Example 8- Design of oligonucleotide probes

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[0357] By way of example, this section describes the design of four oligonucleotide probes using Array Designer Ver 1.1 (Premier Biosoft International, Palo Alto, CA). The major steps in the process are given first.

[0358] Obtain best possible sequence of mRNA from GenBank. If a full-length sequence reference sequence is not available, a partial sequence is used, with preference for the 3' end over the 5' end. When the sequence is known to represent the antisense strand, the reverse complement of the sequence is used for probe design. For sequences

represented in the subtracted leukocyte expression library that have no significant match in GenBank at the time of probe design, our sequence is used.

[0359] Mask low complexity regions and repetitive elements in the sequence using an algorithm such as RepeatMasker.
[0360] Use probe design software, such as Array Designer, version 1.1, to select a sequence of 50 residues with specified physical and chemical properties. The 50 residues nearest the 3' end constitute a search frame. The residues it contains are tested for suitability. If they don't meet the specified criteria, the search frame is moved one residue closer to the 5' end, and the 50 residues it now contains are tested. The process is repeated until a suitable 50-mer is found.
[0361] If no such 50-mer occurs in the sequence, the physical and chemical criteria are adjusted until a suitable 50-mer is found.

[0362] Compare the probe to dbEST, the UniGene cluster set, and the assembled human genome using the BLASTn search tool at NCBI to obtain the pertinent identifying information and to verify that the probe does not have significant similarity to more than one known gene.

Clone 40H12

[0363] Clone 40H12 was sequenced and compared to the nr, dbEST, and UniGene databases at NCBI using the BLAST search tool. The sequence matched accession number NM_002310, a'curated RefSeq project' sequence, see Pruitt et al. (2000) Trends Genet. 16:44-47, encoding leukemia inhibitory factor receptor (LIFR) mRNA with a reported E value of zero. An E value of zero indicates there is, for all practical purposes, no chance that the similarity was random based on the length of the sequence and the composition and size of the database. This sequence, cataloged by accession number NM_002310, is much longer than the sequence of clone 40H12 and has a poly-A tail. This indicated that the sequence cataloged by accession number NM_002310 is the sense strand and a more complete representation of the mRNA than the sequence of clone 40H 12, especially at the 3' end. Accession number "NM_002310" was included in a text file of accession numbers representing sense strand mRNAs, and sequences for the sense strand mRNAs were obtained by uploading a text file containing desired accession numbers as an Entrez search query using the Batch Entrez web interface and saving the results locally as a FASTA file. The following sequence was obtained, and the region of alignment of clone 40H12 is outlined:

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AGTGTTATCAGCACTGATTGGCCATACAAACTGCCCCTTGATCCATCTTGATGGGGAAAATGTTGCAATC AAGATTCGTAATATTTCTGTTTCTGCAAGTAGTGGAACAAATGTAGTTTTTACAACCGAAGATAACATAT TTGGAACCGTTATTTTTGCTGGATATCCACCAGATACTCCTCAACAACTGAATTGTGAGACACATGATTT AAAAGAAATTATATGTAGTTGGAATCCAGGAAGGGTGACAGCGTTGGTGGGCCCACGTGCTACAAGCTAC ACTTTAGTTGAAAGTTTTTCAGGAAAATATGTTAGACTTAAAAGAGCTGAAGCACCTACAAACGAAAGCT ATCAATTATTATTTCAAATGCTTCCAAATCAAGAAATATATAATTTTACTTTGAATGCTCACAATCCGCT GGGTCGATCACAATCAACAATTTTAGTTAATATAACTGAAAAAGTTTATCCCCATACTCCTACTTCATTC AAAGTGAAGGATATTAATTCAACAGCTGTTAAACTTTCTTGGCATTTACCAGGCAACTTTGCAAAGATTA ATTTTTTATGTGAAATTGAAATTAAGAAATCTAATTCAGTACAAGAGCAGCGGAATGTCACAATCAAAGG AGTAGAAAATTCAAGTTATCTTGTTGCTCTGGACAAGTTAAATCCATACACTCTATATACTTTTCGGATT CGTTGTTCTACTGAAACTTTCTGGAAATGGAGCAAATGGAGCAATAAAAAACAACATTTAACAACAGAAG CCAGTCCTTCAAAGGGGCCTGATACTTGGAGAGAGTGGAGTTCTGATGGAAAAAATTTAATAATCTATTG GAAGCCTTTACCCATTAATGAAGCTAATGGAAAAATACTTTCCTACAATGTATCGTGTTCATCAGATGAG GAAACAÇAGTCCCTTTCTGAAATCCCTGATCCTCAGCACAAAGCAGAGATACGACTTGATAAGAATGACT ACATCATCAGCGTAGTGGCTAAAAATTCTGTGGGCTCATCACCACCTTCCAAAATAGCGAGTATGGAAAT TCCAAATGATGTCTCAAAATAGAACAAGTTGTTGGGATGGGAAAGGGGGATTCTCCTCACCTGGCATTAC ACTGGAGAAAAGTTCCCTCAAACAGCACTGAAACTGTAATAGAATCTGATGAGTTTCGACCAGGTATAAG ATATAATTTTTTCCTGTATGGATGCAGAAATCAAGGATATCAATTATTACGCTCCATGATTGGATATATA GAAGAATTGGCTCCCATTGTTGCACCAAATTTTACTGTTGAGGATACTTCTGCAGATTCGATATTAGTAA AATGGGAAGACATTCCTGTGGAAGAACTTAGAGGCTTTTTAAGAGGATATTTGTTTTACTTTGGAAAAGG AGAAAGAGACACCTCAAGATGAGGGTTTTAGAATCAGGTCGTTCTGACATAAAAGTTAAGAATATTACT GACATATCCCAGAAGACACTGAGAATTGCTGATCTTCAAGGTAAAACAAGTTACCACCTGGTCTTGCGAG AATTATTGCCATTCTCATCCCAGTGGCAGTGGCTGTCATTGTTGGAGTGGTGACAAGTATCCTTTGCTAT CGGAAACGAGAATGGATTAAAGAAACCTTCTACCCTGATATTCCAAATCCAGAAAACTGTAAAGCATTAC AGTTTCAAAAGAGTGTCTGTGAGGGAAGCAGTGCTCTTAAAACATTGGAAATGAATCCTTGTACCCCAAA TAATGTTGAGGTTCTGGAAACTCGATCAGCATTTCCTAAAATAGAAGATACAGAAATAATTTCCCCAGTA GCTGAGCGTCCTGAAGATCGCTCTGATGCAGAGCCTGAAAACCATGTGGTTGTGTCCTATTGTCCACCCA TCATTGAGGAAGAAATACCAAACCCAGCCGCAGATGAAGCTGGAGGGGACTGCACAGGTTATTTACATTGA TGTTCAGTCGATGTATCAGCCTCAAGCAAAACCAGAAGAAGAACAAGAAAATGACCCTGTAGGAGGGGCA GGCTATAAGCCACAGATGCACCTCCCCATTAATTCTACTGTGGAAGATATAGCTGCAGAAGAGGACTTAG ATAAAACTGCGGGTTACAGACCTCAGGCCAATGTAAATACATGGAATTTAGTGTCTCCAGACTCTCCTAG ATCCATAGACAGCAACAGTGAGATTGTCTCATTTGGAAGTCCATGCTCCATTAATTCCCGACAATTTTTG ATTCCTCCTAAAGATGAAGACTCTCCTAAATCTAATGGAGGAGGGTGGTCCTTTACAAACTTTTTTCAGA GTTGCTACATCAGCACTGGGCATTCTTGGAGGGATCCTGTGAAGTATTGTTAGGAGGTGAACTTCACTAC ATGTTAAGTTACACTGAAAGTTCATGTGCTTTTAATGTAGTCTAAAAGCCAAAGTATAGTGACTCAGAAT CCTCAATCCACAAAACTCAAGATTGGGAGCTCTTTGTGATCAAGCCAAAGAATTCTCATGTACTCTACCT TCAAGAAGCATTTCAAGGCTAATACCTACTTGTACGTACATGTAAAACAAATCCCGCCGCAACTGTTTTC TGTTCTGTTGTTGTGGGTTTTCTCATATGTATACTTGGTGGAATTGTAAGTGGATTTGCAGGCCAGGGAG AAAATGTCCAAGTAACAGGTGAAGTTTATTTGCCTGACGTTTACTCCTTTCTAGATGAAAACCAAGCACA AGTGACAGCGATTTAGTGTTTTGTTTGATAAAGTATGCTTATTTCTGTGCCTACTGTATAATGGTTATCA AACAGTTGTCTCAGGGGTACAAACTTTGAAAACAAGTGTGACACTGACCAGCCCAAATCATAATCATGTT GTTGGTTGCCCTAATATTTAAAATTTACACTTCTAAGACTAGAGACCCACATTTTTTAAAAATCATTTTA TTTTGTGATACAGTGACAGCTTTATATGAGCAAATTCAATATTATTCATAAGCATGTAATTCCAGTGACT TACTATGTGAGATGACTACTAAGCAATATCTAGCAGCGTTAGTTCCATATAGTTCTGATTGGATTTCGTT CCTCCTGAGGAGACCATGCCGTTGAGCTTGGCTACCCAGGCAGTGGTGATCTTTGACACCTTCTGGTGGA TGTTCCTCCCACTCATGAGTCTTTTCATCATGCCACATTATCTGATCCAGTCCTCACATTTTTAAATATA AAACTAAAGAGAGAATGCTTCTTACAGGAACAGTTACCCAAGGGCTGTTTCTTAGTAACTGTCATAAACT CCTTCAGCACAGCATCCTCTGCCCACCCTTGTTTCTCATAAGCGATGTCTGGAGTGATTGTGGTTCTTGG AAAAGCAGAAGGAAAAACTAAAAAGTGTATCTTGTATTTTCCCTGCCCTCAGGTTGCCTATGTATTTTAC TTTTTTGGTTGGTTGTTTTTTTTTTCATCTGAGATTCTGTAATGTATTTGCAAATAATGGATCAATT AATTTTTTTGAAGCTCATATTGTATCTTTTTAAAAACCATGTTGTGGAAAAAAGCCAGAGTGACAAGTG ACAAAATCTATTTAGGAACTCTGTGTATGAATCCTGATTTTAACTGCTAGGATTCAGCTAAATTTCTGAG

The FASTA file, including the sequence of NM_002310, was masked using the RepeatMasker web interface (Smit, AFA & Green, P RepeatMasker at http://ftp.genome.washington.edu/RM/RepeatMasker.html, Smit and Green). Specifically, during masking, the following types of sequences were replaced with "N's": SINE/MIR & LINE/L2, LINE/L1, LTR/MaLR, LTR/Retroviral, Alu, and

other low informational content sequences such as simple repeats. Below is the sequence following masking:

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CTCTCTCCCAGAACGTGTCTCTGCAAGGCACCGGGCCCTTTCGCTCTGCAGAACTGCACTTGCAAG GACTGCATTGCACAGATGATGGATATTTACGTATGTTTGAAACGACCATCCTGGATGGTGGACAATAAA AGAATGAGGACTGCTTCAAATTTCCAGTGGCTGTTATCAACATTTATTCTTCTATATCTAATGAATCAA TGTTCTTGGAAAGCACCCTCTGGAACAGGCCGTGGTACTGATTATGAAGTTTGCATTGAAAACAGGTCC CGTTCTTGTTATCAGTTGGAGAAAACCAGTATTAAAATTCCAGCTCTTTCACATGGTGATTATGAAATA ACAATAAATTCTCTACATGATTTTGGAAGTTCTACAAGTAAATTCACACTAAATGAACAAAACGTTTCC TTAATTCCAGATACTCCAGAGATCTTGAATTTGTCTGCTGATTTCTCAACCTCTACATTATACCTAAAG TGGAACGACAGGGGTTCAGTTTTTCCACACCGCTCAAATGTTATCTGGGAAATTAAAGTTCTACGTAAA GAGAGTATGGAGCTCGTAAAATTAGTGACCCACAACACACTCTGAATGGCAAAGATACACTTCATCAC AATCTTCATTTTTCTGGTCTCGAAGAGTGGAGTGACTGGAGCCCTGTGAAGAACATTTCTTGGATACCT GATTCTCAGACTAAGGTTTTTCCTCAAGATAAAGTGATACTTGTAGGCTCAGACATAACATTTTGTTGT GTGAGTCAAGAAAAGTGTTATCAGCACTGATTGGCCATACAAACTGCCCCTTGATCCATCTTGATGGG GAAAATGTTGCAATCAAGATTCGTAATATTTCTGTTTCTGCAAGTAGTGGAACAAATGTTAGTTTTTACA ACCGAAGATAACATATTTGGAACCGTTATTTTTGCTGGATATCCACCAGATACTCCTCAACAACTGAAT TGTGAGACACATGATTTAAAAGAAATTATATGTAGTTGGAATCCAGGAAGGGTGACAGCGTTGGTGGGC CCACGTGCTACAAGCTACACTTTAGTTGAAAGTTTTTCAGGAAAATATGTTAGACTTAAAAGAGCTGAA GCACCTACAAACGAAAGCTATCAATTATTATTTCAAATGCTTCCAAATCAAGAAATATATAAATTTTACT TTGAATGCTCACAATCCGCTGGGTCGATCACAATCAACAATTTTAGTTAATAACTGAAAAAGTTTAT CCCCATACTCCTACTTCATACAAGTGAAGGATATTAATTCAACAGCTGTTAAACTTTCTTGGCATTTA CCAGGCAACTTTGCAAAGATTAATTTTTTATGTGAAATTGAAATTAAGAAATCTAATTCAGTACAAGAG CAGCGGAATGTCACAATCAAAGGAGTAGAAAATTCAAGTTATCTTGTTGCTCTGGACAAGTTAAATCCA TACACTCTATATACTTTTCGGATTCGTTGTTCTACTGAAACTTTCTGGAAATGGAGCAAATGGAGCAAT AAAAAACAACATTTAACAACAGAAGCCAGTCCTTCAAAGGGGCCTGATACTTGGAGAGAGTGGAGTTCT GATGGAAAAATTTAATAATCTATTGGAAGCCTTTACCCATTAATGAAGCTAATGGAAAAATACTTTCC TACAATGTATCGTGTTCATCAGATGAGGAAACACAGTCCCTTTCTGAAATCCCTGATCCTCAGCACAAA GCAGAGATACGACTTGATAAGAATGACTACATCATCAGCGTAGTGGCTAAAAATTCTGTGGGCTCATCA CCACCTTCCAAAATAGCGAGTATGGAAATTCCAAATGATGATCTCAAAATAGAACAAGTTGTTGGGATG GGAAAGGGGATTCTCCTCACCTGGCATTACGACCCCAACATGACTTGCGACTACGTCATTAAGTGGTGT AACTCGTCTCGGTCGGAACCATGCCTTATGGACTGGAGAAAGTTCCCTCAAACAGCACTGAAACTGTA

ATAGAATCTGATGAGTTTCGACCAGGTATAAGATATAATTTTTTCCTGTATGGATGCAGAAATCAAGGA
${\tt TATCAATTATTACGCTCCATGATTGGATATATAGAAGAATTGGCTCCCATTGTTGCACCAAATTTTACTCCCCATTGTTGCACCAAATTTTACTCCCCCATTGTTGCACCAAATTTTACTCCCCCCCATTGTTGCACCAAATTTTACTCCCCCCCC$
$\tt GTTGAGGATACTTCTGCAGATTCGATATTAGTAAAATGGGAAGACATTCCTGTGGAAGAACTTAGAGGGCAGAACTTAGAGGGCAGAACTTAGAGGGCAGAACTTAGAGGGCAGAACTTAGAGGGCAGAACTTAGAGGGCAGAACTTAGAGGGCAGAACTTAGAGGGCAGAACTTAGAGGGCAGAACTTAGAGGGCAGAACTTAGAGGGCAAGAACTTAGAGGGCAAGAACTTAGAGGGCAAGAACTTAGAGGGCAAGAACTTAGAGGGCAAGAACTTAGAGGGCAAGAACTTAGAGGGCAAGAACTTAGAGGGCAAGAACTTAGAGGGCAAGAACTTAGAGGGCAAGAACTTAGAGGGCAAGAACTTAGAGGGCAAGAACTTAGAGGGCAAGAACTTAGAGGGCAAGAACTTAGAGGGCAAGAACTTAGAGGGCAAGAACTTAGAGGGCAAGAACTTAGAGGGCAAGAACTTAGAGGGCAAGAACTTAGAGGGCAAGAACTTAGAGGGCAAGAACTTAGAGGGCAAGAACTTAGAGGGCAAGAACTTAGAGGGCAAGAACTTAGAGAGAACTTAGAGAGAACTTAGAGAGAACTTAGAGAGAACTTAGAGAGAACTTAGAGAGAACTTAGAGAGAACTTAGAGAGAACTTAGAGAGAACTTAGAGAGAACTTAGAGAGAACTTAGAGAGAACTTAGAGAGAACTTAGAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACTAGAGAACAACAGAACAACAACAACAACAACAACAACAACA$
$\tt TTTTTAAGAGGATATTTGTTTTACTTTGGAAAAGGAGAAAGAGACACATCTAAGATGAGGGTTTTAGAAGAGAGAG$
${\tt TCAGGTCGTTCTGACATAAAAGTTAAGAATATTACTGACATATCCCAGAAGACACTGAGAATTGCTGATATCCCAGAAGACACTGAGAATTGCTGATATCCCAGAAGACACTGAGAATTGCTGATATCCCAGAAGACACTGAGAATTGCTGATATCCCAGAAGACACTGAGAATTGCTGATATCCCAGAAGACACTGAGAATTGCTGATATCCCAGAAGACACTGAGAATTGCTGATATCCCAGAAGACACTGAGAATTGCTGATATCCCAGAAGACACTGAGAATTGCTGATATCCCAGAAGACACTGAGAATTGCTGATATCCCAGAAGACACTGAGAATTGCTGATATCCCAGAAGACACTGAGAATTGCTGATATCCCAGAAGACACTGAGAATTGCTGATATCCCAGAAGACACTGAGAATTGCTGATATCCCAGAAGACACTGAGAATTGCTGATATCCCAGAAGACACTGAGAATTGCTGATATCCCAGAAGACACTGAGAATTGCTGATATCCCAGAAGACACTGAGAATTGCTGATATCCCAGAAGACACTGAGAATTGCTGATATCCCAGAAGACACTGAGAATTGCTGATATCCCAGAAGACACTGAGAATTGCTGATATCCCAGAAGACACTGAGAATTGCTGATATCCAGAAGACACTGAGAATTGCTGATATCCAGAAGACACTGAGAATTGCTGATATCAGAATTATCAGAATATTACTGACATATCCCAGAAGAACACTGAGAATTGCTGATATCAGAATTATCAGAATATCAGAATATTACTGACATATCAGAATATTACTGACATATCCCAGAAGAACACACTGAGAATTATCAGAATATTACTGACATATATCAGAATATTACTGACAATATCAGAATATTACTGACAATATATCAGAATATTACTGACAATATTACTGACAATATTACTGACAATATTACTGACAATATTACTGACAATATTACTGACAATATTACAATATTACTGAATATTACTGACAATATTACAATATATAT$
$\tt CTTCAAGGTAAAACAAGTTACCACCTGGTCTTGCGAGCCTATACAGATGGTGGAGTGGGCCCGGAGAAGGTTCAAGGTAAAACAAGTTACCACCTGGTCTTGCGAGCCTATACAGATGGTGGAGTGGGCCCGGAGAAGGTTACAAGATGGTGGAGTGGAGTAAAAAAGAAGGTAAAAAAGAAG$
${\tt AGTATGTATGTGGTGACAAAGGAAAATTCTGTGGGATTAATTA$
${\tt GCTGTCATTGTTGGAGTGGTGACAAGTATCCTTTGCTATCGGAAACGAGAATGGATTAAAGAAACCTTCGGAAACGAGAATGGATTAAAGAAACCTTCGGAAACGAGAATGGATTAAAGAAACCTTCGGAAACGAGAATGGATTAAAGAAACCTTCGGAAACGAGAATGGATTAAAGAAACCTTCGGAAACGAGAATGGATTAAAGAAACCTTCGGAAACGAGAATGGATTAAAGAAACCTTCGGAAACGAGAATGGATTAAAGAAACCTTCGGAAACGAGAATGGATTAAAGAAACCTTCGGAAACGAGAATGGATTAAAGAAACCTTCGGAAACGAGAATGGATTAAAGAAACCTTCGGAAACGAGAATGGATTAAAGAAACCTTCGGAAACGAGAATGGATTAAAGAAACCTTCGGAAACGAGAATGGATTAAAGAAACCTTCGGAAACGAGAATGGATTAAAGAAACCTTCGGAAACGAGAATGGATTAAAGAAACCTTCGGAAACGAGAATGGATTAAAGAAACCTTCGGAAACGAGAATGGATTAAAGAAACCTTCGGAAACGAGAATGGATTAAAAGAAACCTTCGGAAACGAGAATGGATTAAAAGAAACCTTCGGAAACGAGAATGGATTAAAAGAAACCTTCGGAAACGAGAATGGATTAAAAGAAAACCTTCGGAAACGAGAATGGAATGGATTAAAAGAAACCTTCGGAAAACGAGAATGGATTAAAAGAAAACCTTCGGAAAACGAGAAACGAGAATGGATTAAAAGAAAACCTTCGGAAAACGAGAATGGATTAAAAGAAAACCTTCGGAAAACGAGAATGGATGAATGGATGAATGGATGAATGGAATGGATGAATGGATGAATGGATGAATAAT$
${\tt TACCCTGATATTCCAAATCCAGAAAACTGTAAAGCATTACAGTTTCAAAAGAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAGAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAAGAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAGAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAAGAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAAGAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAAGAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAAGAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAAGAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAAGAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAAGAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAAGAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAAGAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAAGAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAAGAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAAGAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAAGAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAAGAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAAAAGAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAAGAGAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAAAGAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAAAGAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAAAGAGAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAAAAGAAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAAAGAGTGTCTGTGAGGGAAGCCATTACAGTTTCAAAAAAGAGTGTCTGTGAGGAAGAGCATTACAGTTTACAGTTTTCAAAAAAGAAGAGTGTCTGTGAGAGAAGAGTGTCTGTGAGAGAGA$
${\tt AGTGCTCTTAAAACATTGGAAATGAATCCTTGTACCCCAAATAATGTTGAGGTTCTGGAAACTCGATCACCCCCAAATAATGTTGAGGTTCTGGAAACTCGATCACCCCAAATAATGTTGAGGTTCTGGAAACTCGATCACCCCAAATAATGTTGAGGTTCTGGAAACTCGATCACCCCAAATAATGTTGAGGTTCTGGAAACTCGATCACCCCAAATAATGTTGAGGTTCTGGAAACTCGATCACCCCAAATAATGTTGAGGTTCTGGAAACTCGATCACCCCAAATAATGTTGAGGTTCTGGAAACTCGATCACCCCAAATAATGTTGAGGTTCTGGAAACTCGATCACCCCAAATAATGTTGAGGTTCTGGAAACTCGATCACCCCAAATAATGTTGAGGTTCTGGAAACTCGATCACACCCCAAATAATGTTGAGGTTCTGGAAACTCGATCACACCACACCACACACA$
${\tt GCATTTCCTAAAATAGAAGATACAGAAATAATTTCCCCAGTAGCTGAGCGTCCTGAAGATCGCTCTGATGCTGAGAGATCGCTCTGATGCTGAGAGATCGCTCTGATGCTGAGAGATCGCTCTGATGCTGAGAGATCGCTCTGATGCTGAGAGATCGCTCTGATGCTGAGAGATCGCTCTGATGATGCTGAGAGATCGCTCTGATGCTGAGAGATCGCTCTGATGCTGAGAGATCGCTCTGATGCTGAGAGATCGCTCTGATGCTGAGAGATCGCTCTGATGCTGAGAGATCGCTCTGATGCTGAGAGATCGCTCTGATGCTGAGAGATCGCTCTGATGCTGAGAGATCGCTCTGATGCTGAGAGATCGCTCTGATGATGCTGAGAGATCGCTCTGATGCTGAGATCGCTCTGATGCTGAGATCGCTCTGATGCTGAGATGCTCTGATGATGATGATGATGATGATGATGATGATGATGATGAT$
${\tt GCAGAGCCTGAAAACCATGTGTTGTCCTATTGTCCACCCATCATTGAGGAAGAAATACCAAACCCACCC$
${\tt GCCGCAGATGAAGCTGGAGGGACTGCACAGGTTATTTACATTGATGTTCAGTCGATGTATCAGCCTCAAGCCAAGCCTCAAGCCTCAAGCCTCAAGCCTCAAGCCTCAAGCCTCAAGCCTCAAGCCTCAAGCCTCAAGCCTCAAGCCTCAAGCCTCAAGCCTCAAGCCTCAAGCCTCAAGCCTCAAGCCTCAAGCCTCAAGCCAAGCCTCAAGCCTCAAGCCTCAAGCCTCAAGCCTCAAGCCTCAAGCCTCAAGCCTCAAGCCAAGCAAG$
GCAAAACCAGAAGAAGAACAAGAAAATGACCCTGTAGGAGGGCAGGCTATAAGCCACAGATGCACCTC
$\tt CCCATTAATTCTACTGTGGAAGATATAGCTGCAGAAGAGGACTTAGATAAAACTGCGGGTTACAGACCTGCAGAAGAGAGAG$
${\tt CAGGCCAATGTAAATACATGGAATTTAGTGTCTCCAGACTCTCCTAGATCCATAGACAGCAACAGTGAGCAGCAACAGTGAGCAACAGTGAGCAACAGTGAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGTGAGACAGCAACAGCAACAGTGAGACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAGCAACAA$
${\tt ATTGTCTCATTTGGAAGTCCATGCTCCATTAATTCCCGACAATTTTTGATTCCTCCTAAAGATGAAGACCATGCTCCATTAATTCCCGACAATTTTTGATTCCTCCTAAAGATGAAGACCATGCATG$
${\tt TCTCCTAAATCTAATGGAGGAGGGTGGTCCTTTACAAACTTTTTCAGAACAAACCAAACGATTAACAGGAGGAGGAGGAGGAGGAGGAGGAGGAGGAGGAGGA$
${\tt TGTCACCGTGTCACTCAGTCAGCCATCTCAATAAGCTCTTACTGCTAGTGTTGCTACATCAGCACTGGCTGTGCTACTGCTAGTGTTGCTACATCAGCACTGGGTGTGTTGCTACATCAGCACTGGGTGTGTTGCTACATCAGCACTGGGTGTGTTGCTACATCAGCACTGGGTGTGTTGCTACATCAGCACTGGGTGTGTTGCTACATCAGCACTGGGTGTGTTGCTACATCAGCACTGGGTGTGTTGCTACATCAGCACTGGGTGTGTGT$
${\tt GCATTCTTGGAGGGATCCTGTGAAGTATTGTTAGGAGGTGAACTTCACTACATGTTAAGTTACACTGAAGTTAGGAGGAGGTGAACTTCACTACATGTTAAGTTACACTGAAGTTAGGAGGAGGTGAACTTCACTACATGTTAAGTTACACTGAAGTTAGGAGGAGGTGAACTTCACTACATGTTAAGTTACACTGAAGTTAGGAGGTGAACTTCACTACATGTTAAGTTACACTGAAGTTAGGAGGTGAACTTCACTACATGTTAAGTTACACTGAAGTTACACTGAAGTTAGGAGGTGAACTTCACTACATGTTAAGGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTAGGAGGTGAACTTCACTACATGTTAAGGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTTACACTGAAGTACACTGAAGTACACTGAAGTACACTGAAGTACACTGAAGTACACTGAAGTACACTGAAGTACACTACACTGAAGTACACTGAAGTACACACTGAAGAAGTACACTACACACTACACACTACACACAC$
${\tt AGTTCATGTGCTTTTAATGTAGTCTAAAAGCCAAAGTATAGTGACTCAGAATCCTCAATCCACAAAACTCAGAATCCTCAATCCACAAAACTCAGAATCCTCAATCCACAAAACTCAGAAACTCAGAATCCACAAAACTCAGAATCCACAAAACTCAGAATCCACAAAACTCAGAATCCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAACTCACAAAAAA$
${\tt CAAGATTGGGAGCTCTTTGTGATCAAGCCAAAGAATTCTCATGTACTCTACCTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCAAGAAGTACTCTACCTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGCATTTCAAGAAGAAGCATTTCAAGAAGAAGCATTTCAAGAAGAAGCATTTCAAGAAGAAGCATTTCAAGAAGAAGCATTTCAAGAAGAAGAAGCATTTCAAGAAGAAGAAGAAGAAGAAGAAGAAGAAGAAGAAGAAG$
${\tt GGCTAATACCTACTTGTACGTACATGTAAAACAAATCCCGCCGCAACTGTTTTCTGTTCTGTTTGT}$
${\tt GGTTTTCTCATATGTATACTTGGTGGAATTGTAAGTGGATTTGCAGGCCAGGGAGAAAATGTCCAAGTAGTGTGGAGGAGAAAATGTCCAAGTAGTGGAGGAGAAAATGTCCAAGTAGTGGAGGAGAAAATGTCCAAGTAGTGGAGGAGAAAATGTCCAAGTAGTGGAGGAGAAAATGTCCAAGTAGTGGAGGAGAAAATGTCCAAGTAGTGGAGGAGAAAATGTCCAAGTAGTGGAGGAGAAAATGTCCAAGTAGTGGAGGAGAAAATGTCCAAGTAGTGGAGGAGAAAATGTCCAAGTAGTGGAGGAGAAAATGTCCAAGTAGTGGAGGAGAAAATGTCCAAGTAGTGGAGGAGAAAATGTCCAAGTAGTGGAGGAGAAAAATGTCCAAGTAGTAGTAGTGGAGGAGAAAATGTCCAAGTAGTAGTAGTAGTAGTAGTAGTAGTAGTAGTAGTAGT$
ACAGGTGAAGTTTATTTGCCTGACGTTTACTCCTTTCTAGATGAAAACCAAGCACAGATTTTAAAACTT
CTAAGATTATTCTCCTCTATCCACAGCATTCACNNNNNNNNNN
TTAGTGTTTTGTTTGATAAAGTATGCTTATTTCTGTGCCTACTGTATAATGGTTATCAAACAGTTGTCT
CAGGGGTACAAACTTTGAAAACAAGTGTGACACTGACCAGCCCAAATCATATCATGTTTTCTTGCTGT
GATAGGTTTTGCTTGCCTTTTCATTATTTTTTAGCTTTATGCTTGCT
CCTAATATTTAAAATTTACACTTCTAAGACTAGAGACCCACATTTTTTAAAAATCATTTTATTTTGTGA
TACAGTGACAGCTTTATATGAGCAAATTCAATATTATTCATAAGCATGTAATTCCAGTGACTTACTATG
TGAGATGACTACTAAGCAATATCTAGCAGCGTTAGTTCCATATAGTTCTGATTGGATTTCGTTCCTCCT
GAGGAGACCATGCCGTTGAGCTTGGCTACCCAGGCAGTGGTGATCTTTGACACCTTCTGGTGGATGTTC
CTCCCACTCATGAGTCTTTTCATCATGCCACATTATCTGATCCAGTCCTCACATTTTTAAATATAAAAC
TAAAGAGAGAATGCTTCTTACAGGAACAGTTACCCAAGGGCTGTTTCTTAGTAACTGATAAACTGAT
CTGGATCCATGGGCATACCTGTGTTCGAGGTGCAGCAATTGCTTGGTGAGCTGTGCAGAATTGATTG
TTCAGCACAGCATCCTCTGCCCACCCTTGTTTCTCATAAGCGATGTCTGGAGTGATTGTGGTTCTTGGA
AAAGCAGAAGGAAAAACTAAAAAGTGTATCTTGTATTTTCCCTGCCCTCAGGTTGCCTATGTATTTTAC
CTTTTCATATTTAAGGCAAAAGTACTTGAAAATTTTAAGTGTCCGAATAAGATATGTCTTTTTTGTTTG
TTTTTTTTGGTTGGTTGTTTTTTTTTTTTTTTTTGCAAATAATGCAAATAATGGATCAA

[0364] The length of this sequence was determined using batch, automated computational methods and the sequence, as sense strand, its length, and the desired location of the probe sequence near the 3' end of the mRNA was submitted to Array Designer Ver 1.1 (Premier Biosoft International, Palo Alto, CA). Search quality was set at 100%, number of best probes set at 1, length range set at 50 base pairs, Target Tm set at 75 C. degrees plus or minus 5 degrees, Hairpin max deltaG at 6.0 -kcal/mol., Self dimmer max deltaG at 6.0 -kcal/mol, Run/repeat (dinucleotide) max length set at 5, and Probe site minimum overlap set at 1. When none of the 49 possible probes met the criteria, the probe site would be moved 50 base pairs closer to the 5' end of the sequence and resubmitted to Array Designer for analysis. When no possible probes met the criteria, the variation on melting temperature was raised to plus and minus 8 degrees and the number of identical basepairs in a run increased to 6 so that a probe sequence was produced.

[0365] In the sequence above, using the criteria noted above, Array Designer Ver 1.1 designed a probe corresponding to oligonucleotide number 3037 and is indicated by underlining in the sequence above. It has a melting temperature of 68.4 degrees Celsius and a max run of 6 nucleotides and represents one of the cases where the criteria for probe design in Array Designer Ver 1.1 were relaxed in order to obtain an oligonucleotide near the 3' end of the mRNA (Low melting temperature was allowed).

Clone 463D12

[0366] Clone 463D12 was sequenced and compared to the nr, dbEST, and UniGene databases at NCBI using the BLAST search tool. The sequence matched accession number Al184553, an EST sequence with the definition line "qd60a05.x1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1733840 3' similar to gb:M29550 PROTEIN PHOS-PHATASE 2B CATALYTIC SUBUNIT I (HUMAN);, mRNA sequence." The E value of the alignment was 1.00×10^{-118} . The GenBank sequence begins with a poly-T region, suggesting that it is the antisense strand, read 5' to 3'. The beginning of this sequence is complementary to the 3' end of the mRNA sense strand. The accession number for this sequence was included in a text file of accession numbers representing antisense sequences. Sequences for antisense strand mRNAs were obtained by uploading a text file containing desired accession numbers as an Entrez search query using the Batch Entrez web interface and saving the results locally as a FASTA file. The following sequence was obtained, and the region of alignment of clone 463D12 is outlined:

The FASTA file, including the sequence of AA184553, was then masked using the RepeatMasker web interface, as

shown below. The region of alignment of clone 463D12 is outlined.

[0367] The sequence was submitted to Array Designer as described above, however, the desired location of the probe was indicated at base pair 50 and if no probe met the criteria, moved in the 3' direction. The complementary sequence from Array Designer was used, because the original sequence was antisense. The oligonucleotide designed by Array Designer corresponds to oligonucleotide number 3054 and is complementary to the underlined sequence above. The probe has a melting temperature of 72.7 degrees centigrade and a max run of 4 nucleotides.

Clone 72D4

[0368] Clone 72D4 was sequenced and compared to the nr, dbEST, and UniGene databases at NCBI using the BLAST search tool. No significant matches were found in any of these databases. When compared to the human genome draft, significant alignments were found to three consecutive regions of the reference sequence NT_008060, as depicted below, suggesting that the insert contains three spliced exons of an unidentified gene.

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[0369] Because the reference sequence contains introns and may represent either the coding or noncoding strand for this gene, BioCardia's own sequence file was used to design the oligonucleotide. Two complementary probes were designed to ensure that the sense strand was represented. The sequence of the insert in clone 72D4 is shown below, with the three putative exons outlined.

[0370] The sequence was submitted to RepeatMasker, but no repetitive sequences were found The sequence shown above was used to design the two 50-mer probes using Array Designer as described above. The probes are shown in bold typeface in the sequence depicted below. The probe in the sequence is oligonucleotide number 3020 (SEQ ID NO: 3020) and the complementary probe is oligonucleotide number 318 (SEQ ID NO:318). A portion of the target sequence is listed below (SEQ ID: 3106).

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CAGGTCACACAGCACATCAGTGGCTACATGTGAGCTCAGACCTGGGTCTGCTGCTGTCTTCCCAA
TATCCATGACCTTGACTGATGCAGGTGTCTAGGGATACGTCCATCCCCGTCCTGCTGGAGCCCAGAGCA
CGGAAGCCTGGCCCTCCGAGGAGACAGAAGGGAGTGTCGGACACCATGACGAGAGCTTGGCAGAATAAA
TAACTTCTTTAAACAATTTTACGGCATGAAGAAATCTGGACCAGTTTATTAAATGGGATTTCTGCCACA
AACCTTGGAAGAATCACATCATCTTANNCCCAAGTGAAAACTGTGTTGCGTAACAAAGAACATGACTGC
GCTCCACACATACATCATTGCCCGGCGAGGCGGGACACAAGTCAACGACGGAACACTTGAGACAGGCCT
ACAACTGTGCACGGGTCAGAAGCAAGTTTAAGCCATACTTGCTGCAGTGAGACTACATTTCTGTCTATA

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GAAGATACCTGACTTGATCTGTTTTTCAGCTCCAGTTCCCAGATGTGC

←----GTCAAGGGTCTACACG

 ${\tt GTGTTGTGGTCCCCAAGTATCACCTTCCAATTTCTGGGAG--} {\rightarrow}$

CACAACACCAGGGGTTCATAGTGGAAGGTTAAAG-5'

 $CAGTGCTCTGGCCGGATCCTTGCCGCGCGGATAAAAACT--- \rightarrow$

45 Confirmation of probe sequence

[0371] Following probe design, each probe sequence was confirmed by comparing the sequence against dbEST, the UniGene cluster set, and the assembled human genome using BLASTn at NCBI. Alignments, accession numbers, gi numbers, UniGene cluster numbers and names were examined and the most common sequence used for the probe.

Example 9 - Production of an array of 8000 spotted 50mer oligonucleotides

[0372] We produced an array of 8000 spotted initial candidate 50mer oligonucleotides. Example 8 exemplifies the design and selection of probes for this array.

[0373] Sigma-Genosys (The Woodlands, TX) synthesized un-modified 50-mer oligonucleotides using standard phosphoramidite chemistry, with a starting scale of synthesis of 0.05 µmole (see, e.g., R. Meyers, ed. (1995) Molecular Biology and Biotechnology: A Comprehensive Desk Reference). Briefly, to begin synthesis, a 3' hydroxyl nucleoside with a dimethoxytrityl (DMT) group at the 5' end was attached to a solid support. The DMT group was removed with

trichloroacetic acid (TCA) in order to free the 5'-hydroxyl for the coupling reaction. Next, tetrazole and a phosphoramidite derivative of the next nucleotide were added. The tetrazole protonates the nitrogen of the phosphoramidite, making it susceptible to nucleophilic attack. The DMT group at the 5'-end of the hydroxyl group blocks further addition of nucleotides in excess. Next, the inter-nucleotide linkage was converted to a phosphotriester bond in an oxidation step using an oxidizing agent and water as the oxygen donor. Excess nucleotides were filtered out and the cycle for the next nucleotide was started by the removal of the DMT protecting group. Following the synthesis, the oligo was cleaved from the solid support. The oligonucleotides were desalted, resuspended in water at a concentration of 100 or 200 μ M, and placed in 96-deep well format. The oligonucleotides were re-arrayed into Whatman Uniplate 384-well polyproylene V bottom plates. The oligonucleotides were diluted to a final concentration 30 μ M in 1X Micro Spotting Solution Plus (Telechem/arrayit.com, Sunnyvale, CA) in a total volume of 15 μ I. In total, 8,031 oligonucleotides were arrayed into twenty-one 384-well plates.

[0374] Arrays were produced on Telechem/arrayit.com Super amine glass substrates (Telechem/arrayit.com), which were manufactured in 0.1 mm filtered clean room with exact dimensions of 25x76x0.96 mm. The arrays were printed using the Virtek Chipwriter with a Telechem 48 pin Micro Spotting Printhead. The Printhead was loaded with 48 Stealth SMP3B TeleChem Micro Spotting Pins, which were used to print oligonucleotides onto the slide with the spot size being 110-115 microns in diameter.

Example 10: Identification of diagnostic nucleotide sets for diagnosis of Cardiac Allograft Rejection

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[0375] Genes were identified which have expression patterns useful for the diagnosis and monitoring of cardiac allograft rejection. Further, sets of genes that work together in a diagnostic algorithm for allograft rejection were identified. Patients, patient clinical data and patient samples used in the discovery of markers below were derived from a clinical study described in example 5.

[0376] The collected clinical data is used to define patient or sample groups for correlation of expression data. Patient groups are identified for comparison, for example, a patient group that possesses a useful or interesting clinical distinction, verses a patient group that does not possess the distinction. Measures of cardiac allograft rejection are derived from the clinical data described above to divide patients (and patient samples) into groups with higher and lower rejection activity over some period of time or at any one point in time. Such data are rejection grade as determined from pathologist reading of the cardiac biopsies and data measuring progression of end-organ damage, including depressed left ventricular dysfunction (decreased cardiac output, decreased ejection fraction, clinical signs of low cardiac output) and usage of inotropic agents (Kobashigawa 1998).

[0377] Expression profiles correlating with occurrence of allograft rejection are identified, including expression profiles corresponding to end-organ damage and progression of end-organ damage. Expression profiles are identified predicting allograft rejection, and response to treatment or likelihood of response to treatment. Subsets of the candidate library (or a previously identified diagnostic nucleotide set) are identified, that have predictive value for the presence of allograft rejection or prediction of allograft rejection or end organ damage.

[0378] Mononuclear RNA samples were collected from patients who had recently undergone a cardiac allograft transplantation using the protocol described in example 2. The allograft rejection status at the time of sample collection was determined by examination of cardiac biopsies as described in example 5.

[0379] 180 samples were included in the analysis. Each patient sample was associated with a biopsy and clinical data collected at the time of the sample. The cardiac biopsies were graded by a pathologist at the local center and by a centralized pathologist who read the biopsy slides from all four local centers in a blinded manner. Biopsy grades included 0, 1A, 1B, 2, 3A, and 3B. No grade 4 rejection was identified. Dependent variables were developed based on these grades using either the local center pathology reading or the higher of the two readings, local or centralized. The dependent variables used for correlation of gene expression profiles with cardiac allograft rejection are shown in Table 4. Dependent variables are used to create classes of samples corresponding to the presence or absence of rejection.

[0380] Clinical data were also used to determine criteria for including samples in the analysis. The strictest inclusion criteria required that samples be from patients who did not have a bacterial or viral infection, were at least two weeks post cardiac transplant and were not currently admitted to the hospital. A second inclusion criteria (inclusion 2) reduced the post-transplant criteria to 1 week and eliminated the hospital admission criteria.

[0381] After preparation of RNA (example 2), amplification, labeling, hybridization, scanning, feature extraction and data processing were done as described in Example 11, using the oligonucleotide microarrays described in Example 9. The resulting log ratio of expression of Cy3 (patient sample)/ Cy5 (R50 reference RNA) was used for analysis. This dataset is called the "static" data. A second type of dataset, referenced, was derived from the first. These datasets compared the gene expression log ratio in each sample to a baseline sample from the same patient using the formula:

ref log ratio =
$$(\log ratio_{sample}) - (\log ratio_{baseline})$$

[0382] Two referenced datasets were used, named "0 HG" and "Best 0". The baseline for 0 HG was a Grade 0 sample from the same patient as the sample, using the highest grade between the centralized and local pathologists. The baseline for Best 0 was a Grade 0 sample from the same patient as the sample, using both the local and centralized reader biopsy grade data. When possible a Grade 0 prior to the sample was used as the baseline in both referenced datasets.

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[0383] The datasets were also divided into subsets to compare analysis between two subsets of roughly half of the data. The types of subsets constructed were as follows. First half/second half subsets were the first half of the samples and the second half of the samples from a dataset ordered by sample number. Odd/even subsets used the same source, a dataset ordered by sample number, but the odd subset consisted of every 2nd sample starting with the first and the even subset consisted of every 2nd sample starting with the second sample, Center 14/other subsets were the same datasets, divided by transplant hospital. The center 14 subset consisted of all samples from patients at center 14, while the other subset consisted of all samples from the other three centers (12,13, and 15).

[0384] Initially, significance analysis for microarrays (SAM, Tusher 2001, Example 15) was used to discover genes that were differentially expressed between the rejection and no-rejection groups. Ninety-six different combinations of dependent variables, inclusion criteria, static/referenced, and data subsets were used in SAM analysis to develop the primary lists of genes significantly differentially expressed between rejection and no-rejection. The most significant of these genes were chosen based on the following criteria. Tier 1 genes were those which appeared with an FDR of less than 20% in identical analyses in two independent subsets. Tier 2 genes were those which appeared in the top 20 genes on the list with an FDR less than 20% more than 50% of the time over all dependent variables with the inclusion criteria, and static/referenced constant. Tier 3 genes were those that appeared more than 50% of the time with an FDR less than 20% more than 50% of the time over all dependent variables with the inclusion criteria, and static/referenced constant. The genes that were identified by the analysis as statistically differentially expressed between rejection and no rejection are shown in Table 2.

[0385] SAM chooses genes as significantly different based on the magnitude of the difference between the groups and the variation among the samples within each group. An example of the difference between some Grade 0 and some Grade 3A samples for 9 genes is shown in Figure 7A.

[0386] Additionally, many of these same combinations were used in the Supervised Harvesting of Expression Trees (SHET, Hastie et al. 2001) algorithm (see example 15) to identify markers that the algorithm chose as the best to distinguish between the rejection and no rejection classes using a bias factor of 0.01. The top 20 or 30 terms were taken from the SHET output and among all comparisons in either the static or referenced data the results were grouped. Any gene found in the top 5 terms in more than 50% of the analyses was selected to be in group B1 (Table 2). The occurrences of each gene were tabulated over all SHET analysis (for either static or referenced data) and the 10 genes that occurred the most were selected to be in group B2 (Table 2).

[0387] An additional classification method used was CART (Salford Systems, San Diego, example 15). Either the static or referenced dataset was reduced to only the genes for which expression values (log ratios) were present in at least 80% of the samples. These data were used in CART with the default settings, using the Symmetric Gini algorithm. Each of the dependent variables was used with both the full sample set and the strict inclusion criteria. Two groups of genes were identified. Group C1 were those genes that were a primary splitter (1st decision node). Group C2 genes were the 10 genes that occurred as splitters the most often over all these analyses.

[0388] Two other classification models were developed and their best genes identified as markers of cardiac allograft rejection. Group D genes were identified from a set of 59 samples, referenced data, local biopsy reading grade, using logistic regression. Group E genes were identified from the primary static dataset using a K-nearest neighbor classification algorithm.

[0389] Both hierarchical clustering (Eisen et al. 1998) and CART were used to identify surrogates for each identified marker. Hierarchical clustering surrogates are genes co-expressed in these and were chosen from the nearest branches of the dendrogram. CART surrogates were identified by CART as the surrogates for those genes chosen as primary splitters at decision nodes.

[0390] Primers for real-time PCR validation were designed for each of the marker genes as described in Example 13. [0391] CART was used to build a decision tree for classification of samples as rejection or no-rejection using the gene expression data from the arrays. The analysis identified sets of genes that can be used together to accurately identify samples derived from cardiac allograft transplant patients. The set of genes and the identified threshold expression levels for the decision tree are referred to as a "models". This model can be used to predict the rejection state of an unknown sample. The input data were the static expression data (log ratio) and the referenced expression data (log ratio referenced to the best available grade 0 from either the centralized reader or the local reader) for 139 of our top

marker genes. These two types of expression data were entered into the CART software as independent variables. The dependent variable was rejection state, defined for this model as no rejection = grade 0 and rejection = grade 3A. Samples were eliminated from consideration in the training set if they were from patients with either bacterial or viral infection or were from patients who were less than two weeks post-transplant. The method used was Symmetric Gini, allowing linear combinations of independent variables. The costs were set to I for both false negatives and false positives and the priors were set equal for the two states. No penalties were assessed for missing data, however the marker genes selected have strong representation across the dataset. 10-fold cross validation was used to test the model. Settings not specified remained at the default values.

[0392] The model shown in Figure 7B is based on decisions about expression values at three nodes, each a different marker gene. The cost assigned to this model is 0.292, based on the priors being equal, the costs set to 1 for each type of error, and the results from the 10-fold cross validation.

[0393] In the training set, no rejection samples were misclassified (sensitivity = 100%) and only 1 no-rejection sample was misclassified (specificity = 94.4%). Following 10-fold cross validation, 2 rejection samples were misclassified (sensitivity = 87.5%) and 3 no-rejection samples were misclassified (specificity = 83.3%). The CART software assigns surrogate markers for each decision node.

[0394] These genes can be used alone or in association with other genes or variables to build a diagnostic gene set or a classification algorithm. These genes can be used in association with known gene markers for rejection (such as those identified in the prior art) to provide a diagnostic algorithm.

Example 11- Amplification, labeling, and hybridization of total RNA to an oligonucleotide microarray Amplification, labeling, hybridization and scanning

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[0395] Samples consisting of at least 0.5 to 2 μ g of intact total RNA were further processed for array hybridization. When available, 2 μ g of intact total RNA is used for amplification. Amplification and labeling of total RNA samples was performed in three successive enzymatic reactions. First, a single-stranded DNA copy of the RNA was made (hereinafter, "ss-cDNA"). Second, the ss-cDNA was used as a template for the complementary DNA strand, producing double-stranded cDNA (hereinafter, "ds-cDNA, or cDNA"). Third, linear amplification was performed by in vitro transcription from a bacterial T_7 promoter. During this step, fluorescent-conjugated nucleotides were incorporated into the amplified RNA (hereinafter, "aRNA").

[0396] The first strand cDNA was produced using the Invitrogen kit (Superscript II). The first strand cDNA was produced in a reaction composed of 50 mM Tris-HCl (pH 8.3), 75 mM KCl, and 3 mM MgCl₂ (1x First Strand Buffer, Invitrogen), 0.5 mM dGTP, 0.5 mM dATP, 0.5 mM dTTP, 0.5 mM dCTP, 10 mM DTT, 200 U reverse transcriptase (Superscript II, Invitrogen, #18064014), 15 U RNase inhibitor (RNAGuard, Amersham Pharmacia, #27-0815-01), 5 μM T7T24 primer 3105) and 0.5 to 2 μg of selected sample total RNA. Several purified, recombinant control mRNAs from the plant Arabidopsis thaliana were added to the reaction mixture: 2-20 pg of the following genes CAB, RCA, LTP4, NAC1, RCP1,XCP2, RBCL, LTP6, TIM, and PRKase (Stratagene, #252201, #252202, #252204, #252208, #252207, #252206, #252203, #252205, #252209, #252210 respectively). The control RNAs allow the estimate of copy numbers for individual mRNAs in the clinical sample because corresponding sense oligonucleotide probes for each of these plant genes are present on the microarray. The final reaction volume of 20 µl was incubated at 42°C for 90 min. For synthesis of the second cDNA strand, DNA polymerase and RNase were added to the previous reaction, bringing the final volume to 150 µl. The previous contents were diluted and new substrates were added to a final concentration of 20 mM Tris-HCl (pH 7.0) (Fisher Scientific, Pittsburgh, PA #BP1756-100), 90 mMKCI (Teknova, Half Moon Bay, CA, #0313-500), 4.6 mM MgCl₂ (Teknova, Half Moon Bay, CA, #0304-500), 10 mM(NH₄)₂SO₄ (Fisher Scientific #A702-500)(1 x Second Strand buffer, Invitrogen), 0.266 mM dGTP, 0.266 mM dATP, 0.266 mM dTTP, 0.266 mM dCTP, 40 U E. coli DNA polymerase (Invitrogen, #18010-025), and 2 U RNaseH (Invitrogen, #18021-014). The second strand synthesis took place at 16°C for 150 minutes.

[0397] Following second-strand synthesis, the ds-cDNA was purified from the enzymes, dNTPs, and buffers before proceeding to amplification, using phenol-chloroform extraction followed by ethanol precipitation of the cDNA in the presence of glycogen.

[0398] Alternatively, a silica-gel column is used to purify the cDNA (e.g. Qiaquick PCR cleanup from Qiagen, #28104). The volume of the column purified cDNA was reduced by ethanol precipitation in the presence of glycogen in which the cDNA was collected by centrifugation at > 10,000 xg for 30 minutes, the supernatant is aspirated, and 150 μ l of 70% ethanol, 30% water was added to wash the DNA pellet. Following centrifugation, the supernatant was removed, and residual ethanol was evaporated at room temperature. Alternatively, the volume of the column purified cDNA is reduce in a vacuum evaporator where the supernatant is reduce to a final volume of 7.4 μ l.

[0399] Linear amplification of the cDNA was performed by in vitro transcription of the cDNA. The cDNA pellet from the step described above was resuspended in 7.4 μ l of water, and in vitro transcription reaction buffer was added to a

final volume of 20 μ l containing 7.5 mM GTP, 7.5 mM ATP, 7.5 mM TTP, 2.25 mM CTP, 1.025 mM Cy3-conjugated CTP (Perkin Elmer; Boston, MA, #NEL-580), 1 x reaction buffer (Ambion, Megascript Kit, Austin, TX and # 1334) and 1 % T₇ polymerase enzyme mix (Ambion, Megascript Kit, Austin, TX and #1334). This reaction was incubated at 37°C overnight. Following in vitro transcription, the RNA was purified from the enzyme, buffers, and excess NTPs using the RNeasy kit from Qiagen (Valencia, CA; # 74106) as described in the vendor's protocol. A second elution step was performed and the two eluates were combined for a final volume of 60 μ l. RNA is quantified using an Agilent 2100 bioanalyzer with the RNA 6000 nano LabChip.

[0400] Reference RNA was prepared as described above, except Cy5-CTP was incorporated instead of Cy3CTP. Reference RNA from five reactions, each reaction started with 2 ug total RNA, was pooled together and quantitated as described above.

Hybridization to an array

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[0401] RNA was prepared for hybridization as follows: for an $18\text{mm}\times55\text{mm}$ array, $20~\mu g$ of amplified RNA (aRNA) was combined with $20~\mu g$ of reference aRNA. The combined sample and reference aRNA was concentrated by evaporating the water to $10~\mu l$ in a vacuum evaporator. The sample was fragmented by heating the sample at 95°C for 30~minutes to fragment the RNA into 50-200~bp pieces. Alternatively, the combined sample and reference aRNA was concentrated by evaporating the water to $5~\mu l$ in a vacuum evaporator. Five μl of 20~mM zinc acetate was added to the aRNA and the mix incubated at 60°C for 10~minutes. Following fragmentation, $40~\mu l$ of hybridization buffer was added to achieve final concentrations of $5\times\text{SSC}$ and 0.20~MSDS with $0.1~\mu g/u l$ of Cot-1 DNA (Invitrogen) as a competitor DNA. The final hybridization mix was heated to 98°C , and then reduced to 50°C at 0.1°C per second.

[0402] Alternatively, formamide is included in the hybridization mixture to lower the hybridization temperature.

[0403] The hybridization mixture was applied to a pre-heated 65°C microarray, surface, covered with a glass coverslip (Corning, #2935-246), and placed on a pre-heated 65°C hybridization chamber (Telechem, AHC-10). 15 ul of 5xSSC was placed in each of the reservoir in the hybridization chamber and the chamber was sealed and placed in a water bath at 62°C for overnight (16-20 hrs). Following incubation, the slides were washed in 2xSSC, 0.1% SDS for five minutes at 30°C, then in 2xSSC for another five minutes at 30°C, then in 0.2×SSC for two minutes at room temperature. The arrays were spun at 1000xg for 2 minutes to dry them. The dry microarrays are then scanned by methods described above.

[0404] The microarrays were imaged on the Agilent (Palo Alto, CA) scanner G2565AA. The scan settings using the Agilent software were as follows: for the PMT Sensitivity (100% Red and 100% Green); Scan Resolution (10 microns); red and green dye channels; used the default scan region for all slides in the carousel; using the largest scan region; scan date for Instrument ID; and barcode for Slide ID. The full image produced by the Agilent scanner was flipped, rotated, and split into two images (one for each signal channel) using TIFFSplitter (Agilent, Palo Alto, CA). The two channels are the output at 532 nm (Cy3-labeled sample) and 633 nm (Cy5-labeled R50). The individual images were loaded into GenePix 3.0 (Axon Instruments, Union City, CA) for feature extraction, each image was assigned an excitation wavelength corresponding the file opened; Red equals 633 nm and Green equals 532 nm. The setting file (gal) was opened and the grid was laid onto the image so that each spot in the grid overlaped with >50% of the feature. Then the GenePix software was used to find the features without setting minimum threshold value for a feature. For features with low signal intensity, GenePix reports "not found". For all features, the diameter setting was adjusted to include only the feature if necessary.

[0405] The GenePix software determined the median pixel intensity for each feature (F_i) and the median pixel intensity of the local background for each feature (B_i) in both channels. The standard deviation (SDF_i) and SDB_i for each is also determined. Features for which GenePix could not discriminate the feature from the background were "flagged" as described below.

[0406] Following feature extraction into a ".gpr" file, the header information of the .gpr file was changed to carry accurate information into the database. An Excel macro was written to include the following information: Name of the original .tif image file, SlidelD, Version of the feature extraction software, GenePix Array List file, GenePix Settings file, ScanID, Name of person who scanned the slide, Green PMT setting, Red PMT setting, ExtractID (date .gpr file was created, formatted as yyyy.mm.dd-hh.mm.ss), Results file name (same as the .gpr file name), StorageCD, and Extraction comments.

Pre-processing with Excel Templates

[0407] Following analysis of the image and extraction of the data, the data from each hybridization was preprocessed to extract data that was entered into the database and subsequently used for analysis. The complete GPR file produced by the feature extraction in GenePix was imported into an excel file pre-processing template or processed using a AWK script. Both programs used the same processing logic and produce identical results. The same excel template or AWK

script was used to process each GPR file. The template performs a series of calculations on the data to differentiate poor features from others and to combine duplicate or triplicate feature data into a single data point for each probe.

[0408] The data columns used in the pre-processing were: Oligo ID, F633 Median (median value from all the pixels in the feature for the Cy5 dye), B633 Median (the median value of all the pixels in the local background of the selected feature for Cy5), B633 SD (the standard deviation of the values for the pixels in the local background of the selected feature for Cy5), F532 Median (median value from all the pixels in the feature for the Cy3 dye), B532 Median (the median value of all the pixels in the local background of the selected feature for Cy3), B532 SD (the standard deviation of the values for the pixels in the local background of the selected feature for Cy3), and Flags. The GenePix Flags column contains the flags set during feature extraction. "-75" indicates there were no features printed on the array in that position, "-50" indicates that GenePix could not differentiate the feature signal from the local background, and "-100" indicates that the user marked the feature as bad.

[0409] Once imported, the data associated with features with -75 flags was not used. Then the median of B633 SD and B532 SD were calculated over all features with a flag value of "0". The minimum values of B633 Median and B532 Median were identified, considering only those values associated with a flag value of "0". For each feature, the signal to noise ratio (S/N) was calculated for both dyes by taking the fluorescence signal minus the local background (BGSS) and dividing it by the standard deviation of the local background:

$$S/N = \frac{F_i - B_i}{SDB_i}$$

[0410] If the S/N was less than 3, then an adjusted background-subtracted signal was calculated as the fluorescence minus the minimum local background on the slide. An adjusted S/N was then calculated as the adjusted background subtracted signal divided by the median noise over all features for that channel. If the adjusted S/N was greater than three and the original S/N were less than three, a flag of 25 was set for the Cy5 channel, a flag of 23 was set for the Cy3 channel, and if both met these criteria, then a flag of 28 was set. If both the adjusted S/N and the original S/N were less than three, then a flag of 65 was set for Cy5, 63 set for Cy3, and 68 set if both dye channels had an adjusted S/N less than three. All signal to noise calculations, adjusted background-subtracted signal, and adjusted S/N were calculated for each dye channel. If the BGSS value was greater than or equal to 64000, a flag was set to indicate saturation; 55 for Cy5, 53 for Cy3, 58 for both.

[0411] The BGSS used for further calculations was the original BGSS if the original S/N was greater than or equal to three. If the original S/N ratio was less than three and the adjusted S/N ratio was greater than or equal to three, then the adjusted BGSS was used. If the adjusted S/N ratio was less than three, then the adjusted BGSS was used, but with knowledge of the flag status.

[0412] To facilitate comparison among arrays, the Cy3 and Cy5 data were scaled. The log of the ratio of Green/Red was determined for all features. The median log ratio value for good features (Flags 0, 23, 25, 28, 63) was determined. The feature values were scaled using the following formula: Log_Scaled_Feature_Ratio = Log_Feature_Ratio_Median_Log_Ratio.

[0413] The flag setting for each feature was used to determine the expression ratio for each probe, a choice of one, two or three features. If all features had flag settings in the same category (categories=negatives, 0 to 28, 53-58, and 63-68), then the average of the three scaled, anti log feature ratios was calculated If the three features did not have flags in the same category, then the feature or features with the best quality flags were used (0>25>23>28>55>53>58>65>63>68). Features with negative flags were never used. When the best flags were two or three features in the same category, the anti log average was used. If a single feature had a better flag category than the other two then the anti log of that feature ratio was used.

[0414] Once the probe expression ratios were calculated from the one, two, or three features, the log of the scaled, averaged ratios was taken as described below and stored for use in analyzing the data. Whichever features were used to calculate the probe value, the flag from those features was carried forward and stored as the flag value for that probe. 2 different data sets can be used for analysis. Flagged data uses all values, including those with flags. Filtered data sets are created by removing flagged data from the set before analysis.

Example 12: Real-time PCR validation of array expression results

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[0415] Leukocyte microarray gene expression was used to discover expression markers and diagnostic gene sets for clinical outcomes. It is desirable to validate the gene expression results for each gene using a more sensitive and quantitative technology such as real-time PCR. Further, it is possible for the diagnostic nucleotide sets to be implemented as a diagnostic test as a real-time PCR panel. Alternatively, the quantitative information provided by real-time PCR

validation can be used to design a diagnostic test using any alternative quantitative or semi-quantitative gene expression technology. To validate the results of the microarray experiments we used real-time, or kinetic, PCR. In this type of experiment the amplification product is measured during the PCR reaction. This enables the researcher to observe the amplification before any reagent becomes rate limiting for amplification. In kinetic PCR the measurement is of C_T (threshold cycle) or C_P (crossing point). This measurement (C_T = C_P) is the point at which an amplification curve crosses a threshold fluorescence value. The threshold is set to a point within the area where all of the reactions were in their linear phase of amplification. When measuring C_T , a lower C_T value is indicative of a higher amount of starting material since an earlier cycle number means the threshold was crossed more quickly.

[0416] Several fluorescence methodologies are available to measure amplification product in real-time PCR. Taqman (Applied BioSystems, Foster City, CA) uses fluorescence resonance energy transfer (FRET) to inhibit signal from a probe until the probe is degraded by the sequence specific binding and Taq 3' exonuclease activity. Molecular Beacons (Stratagene, La Jolla, CA) also use FRET technology, whereby the fluorescence is measured when a hairpin structure is relaxed by the specific probe binding to the amplified DNA. The third commonly used chemistry is Sybr Green, a DNA-binding dye (Molecular Probes, Eugene, OR). The more amplified product that is produced, the higher the signal. The Sybr Green method is sensitive to non-specific amplification products, increasing the importance of primer design and selection. Other detection chemistries can also been used, such as ethedium bromide or other DNA-binding dyes and many modifications of the fluorescent dye/quencher dye Taqman chemistry.

Sample prep and cDNA synthesis

[0417] The inputs for real time PCR reaction are gene-specific primers, cDNA from specific patient samples, and standard reagents. The cDNA was produced from mononuclear RNA (prepared as in example 2) or whole blood RNA by reverse transcription using Oligo dT primers (Invitrogen, 18418-012) and random hexamers (Invitrogen, 48190-011) at a final concentration of 0.5 ng/μl and 3 ng/μl respectively. For the first strand reaction mix, 0.5 μg of mononuclear total RNA or 2 μg of whole blood RNA and I μl of the Oligo dT/ Random Hexamer Mix, were added to water to a final volume of 11.5 μl. The sample mix was then placed at 70°C for 10 minutes. Following the 70°C incubation, the samples were chilled on ice, spun down, and 88.5 μl of first strand buffer mix dispensed into the reaction tube. The final first strand buffer mix produced final concentrations of 1X first strand buffer (Invitrogen, Y00146, Carlsbad, CA), 10 mM DTT (Invitrogen, Y00147), 0.5 mM dATP (NEB, N0440S, Beverly, MA), 0.5 mM dGTP (NEB, N0442S), 0.5 mM dTTP (NEB, N0443S), 0.5 mM dCTP (NEB, N0441S), 200U of reverse transcriptase (Superscript II, Invitrogen, 18064-014), and 18U of RNase inhibitor (RNAGaurd Amersham Pharmacia, 27-0815-01, Piscataway, NJ). The reaction was incubated at 42°C for 90 minutes. After incubation the enzyme was heat inactivated at 70°C for 15 minutes, 2 U of RNAse H added to the reaction tube, and incubated at 37°C for 20 minutes.

PRIMER DESIGN

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[0418] Two methods were used to design primers. The first was to use the software, Primer Expresstm and recommendations for primer design that are provided with the GeneAmp® 7700 Sequence Detection System supplied by Applied BioSystems (Foster City, CA). The second method used to design primers was the PRIMER3 ver 0.9 program that is available from the Whitehead Research Institute, Cambridge, Massachusetts at the Whitehead Research web site. The program can also be accessed on the World Wide Web at the web site at the Massechusetts Institute of Technology website. Primers and Taqman/hybridization probes were designed as described below using both programs. [0419] The Primer Express literature explains that primers should be designed with a melting temperature between 58 and 60 degrees C. while the Taqman probes should have a melting temperature of 68 to 70 under the salt conditions of the supplied reagents. The salt concentration is fixed in the software. Primers should be between 15 and 30 basepairs long. The primers should produce and amplicon in size between 50 and 150 base pairs, have a C-G content between 20% and 80%, have no more than 4 identical base pairs next to one another, and no more than 2 C's and G's in the last 5 bases of the 3' end. The probe cannot have a G on the 5' end and the strand with the fewest G's should be used for the probe.

[0420] Primer3 has a large number of parameters. The defaults were used for all except for melting temperature and the optimal size of the amplicon was set at 100 bases. One of the most critical is salt concentration as it affects the melting temperature of the probes and primers. In order to produce primers and probes with melting temperatures equivalent to Primer Express, a number of primers and probes designed by Primer Express were examined using PRIMER3. Using a salt concentration of 50 mM these primers had an average melting temperature of 3.7 degrees higher than predicted by Primer Express. In order to design primers and probes with equivalent melting temperatures as Primer Express using PRIMER3, a melting temperature of 62.7 plus/minus 1.0 degree was used in PRIMER3 for primers and 72.7 plus/minus 1.0 degrees for probes with a salt concentration of 50 mM.

[0421] The C source code for Primer3 was downloaded and complied on a Sun Enterprise 250 server using the GCC

complier. The program was then used from the command line using a input file that contained the sequence for which we wanted to design primers and probes along with the input parameters as described by help files that accompany the software. Using scripting it was possible to input a number of sequences and automatically generate a number of possible probes and primers.

[0422] Primers for β -Actin (Beta Actin, Genbank Locus: NM_001101) and β -GUS: glucuronidase, beta, (GUSB, Genbank Locus: NM_000181), two reference genes, were designed using both methods and are shown here as examples:

The first step was to mask out repetitive sequences found in the mRNA sequences using RepeatMasker program that can be accessed at: the web site University of Washington Genome Repeatmasker website. (Smit, A.F.A. & Green, P.).

[0423] The last 500 basepairs on the last 3' end of masked sequence was then submitted to PRIMER3 using the following exemplary input sequences:

PRIMER_SEQUENCE_ID=>GUSB (SEQID 3084)
SEQUENCE=GAAGAGTACCAGAAAAGTCTGCTAGAGCAGTACCATCTGGGTCTGGATCAAAAACGCAGA
AAATATGTGGTTGGAGGAGCTCATTTGGAATTTTGCCGATTTCATGACTGAACAGTCACCGACGAGAGTG
CTGGGGAATAAAAAGGGGATCTTCACTCGGCAGAGACAACCAAAAAGTGCAGCGTTCCTTTTGCGAGAG
AGATACTGGAAGATTGCCAATGAAACCAGGTATCCCCACTCAGTAGCCAAGTCACAATGTTTGGAAAAC
AGCCCGTTTACTTGAGCAAGACTGATACCACCTGCGTGTCCCTTCCTCCCCGAGTCAGGGCGACTTCCA
CAGCAGCAGAACAAGTGCCTCCTGGACTGTTCACGGCAGACCGTTTCTGGCCTGGGTTTTGTGG
TCATCTATTCTAGCAGGGAACACTAAAGGTGGAAATAAAAGATTTTCTATTATGGAAATAAAAGAGTTGG
CATGAAAGTCGCTACTG

[0424] After running PRIMER3, 100 sets of primers and probes were generated for ACTB and GUSB. From this set, nested primers were chosen based on whether both left primers could be paired with both right primers and a single Taqman probe could be used on an insert of the correct size. With more experience we have decided not use the mix and match approach to primer selection and just use several of the top pairs of predicted primers.

[0425] For ACTB this turned out to be:

Forward 75 CACAATGTGGCCGAGGACTT(SEQID 3085), Forward 80 TGTGGCCGAGGACTTTGATT(SEQID 3086), Reverse 178 TGGCTTTTAGGATGGCAAGG(SEQID 3087), and Reverse 168 GGGGGCTTAGTTTGCTTCCT(SEQID 3088).

Upon testing, the F75 and R178 pair worked best.

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[0426] For GUSB the following primers were chosen:

Forward 59 AAGTGCAGCGTTCCTTTGC(SEQID 3089), Forward 65 AGCGTTCCTTTTGCGAGAGA (SEQID 3090), Reverse 158 CGGGCTGTTTTCCAAACATT (SEQID 3091), and Reverse 197 GAAGGGACACGCAGGTGGTA (SEQID 3092).

[0427] No combination of these GUSB pairs worked well.

[0428] In addition to the primer pairs above, Primer Express predicted the following primers for GUSB: Forward 178 TACCACCTGCGTGTCCCTTC (SEQID 3093) and Reverse 242 GAGGCACTTGTTCTGCTGCTG (SEQID 3094). This pair of primers worked to amplify the GUSB mRNA.

[0429] The parameters used to predict these primers in Primer Express were:

Primer Tm: min 58, Max=60, opt 59, max difference=2 degrees

Primer GC: min=20% Max =80% no 3' G/C clamp

Primer: Length: min=9 max=40 opt=20 Amplicon: min Tm=0 max Tm=85

min = 50 bp max = 150 bp

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Probe: Tm 10 degrees > primers, do not begin with a G on 5' end

Other: max base pair repeat = 3 max number of ambiguous residues = 0

secondary structure: max consecutive bp = 4, max total bp = 8

10 Uniqueness: max consecutive match = 9

max % match = 75

max 3' consecutive match = 7

[0430] Granzyme B is a marker of transplant rejection.

For Granzyme B the following sequence (NM_004131) (SEQID 3096) was used as input for Primer3:

For Granzyme B the following primers were chosen for testing:

Forward 81 ACGAGCCTGCACCAAAGTCT (SEQID 3097)

Forward 63 AAACAATGGCATGCCTCCAC (SEQID 3098)

Reverse 178 TCATTACAGCGGGGGCTTAG (SEQID 3099)

Reverse 168 GGGGGCTTAGTTTGCTTCCT (SEQID 3100)

[0431] Testing demonstrated that F81 and R178 worked well.

[0432] Using this approach, primers were designed for all the genes that were shown to have expression patterns that correlated with allograft rejection. These primer pairs are shown in Table 2, and are added to the sequence listing. Primers can be designed from any region of a target gene using this approach.

PRIMER ENDPOINT TESTING

[0433] Primers were first tested to examine whether they would produce the correct size product without non-specific amplification. The standard real-time PCR protocol was used without the Rox and Sybr green dyes. Each primer pair was tested on cDNA made from universal mononuclear leukocyte reference RNA that was produced from 50 individuals as described in Example 3 (R50).

[0434] The PCR reaction consisted of 1X RealTime PCR Buffer (Ambion, Austin, TX), 2mM MgCl2 (Applied BioSystems, B02953), 0.2mM dATP (NEB), 0.2mM dTTP (NEB), 0.2mM dCTP (NEB), 0.2mM dGTP (NEB), .625U AmpliTaq Gold (Applied BioSystems, Foster City, CA), 0.3μ M of each primer to be used (Sigma Genosys, The Woodlands, TX), 5μ l of the R50 reverse-transcription reaction and water to a final volume of 19μ l.

[0435] Following 40 cycles of PCR, 10 microliters of each product was combined with Sybr green at a final dilution of 1:72,000. Melt curves for each PCR product were determined on an ABI 7900 (Applied BioSystems, Foster City, CA), and primer pairs yielding a product with one clean peak were chosen for further analysis. One microliter of the product from these primer pairs was examined by agarose gel electrophoresis on an Agilent Bioanalyzer, DNA1000 chip (Palo Alto, CA). Results for 2 genes are shown in Figure 9. From the primer design and the sequence of the target gene, one can calculate the expected size of the amplified DNA product. Only primer pairs with amplification of the desired product and minimal amplification of contaminants were used for real-time PCR. Primers that produced multiple products of different sizes are likely not specific for the gene of interest and may amplify multiple genes or chromosomal loci.

PRIMER OPTIMIZATION/EFFICIENCY

[0436] Once primers passed the end-point PCR, the primers were tested to determine the efficiency of the reaction

in a real-time PCR reaction. cDNA was synthesized from starting total RNA as described above. A set of 5 serial dilutions of the R50 reverse-transcribed cDNA (as described above) were made in water: 1:10, 1:20, 1:40, 1:80, and 1:160. **[0437]** The Sybr Green real-time PCR reaction was performed using the Taqman PCR Reagent kit (Applied BioSystems, Foster City, CA, N808-0228). A master mix was made that consisted of all reagents except the primes and template. The final concentration of all ingredients in the reaction was 1X Taqman Buffer A (Applied BioSystems), 200 μ M dGTP (Applied BioSystems), 200 μ M dGTP (Applied BioSystems), 200 μ M dGTP (Applied BioSystems), 1:400,000 diluted Sybr Green dye (Molecular Probes), 1.25U AmpliTaq Gold (Applied BioSystems). The PCR master mix was dispensed into two, light-tight tubes. Each β-Actin primer F75 and R178 (Sigma-Genosys, The Woodlands, TX), was added to one tube of PCR master mix and Each β-GUS primer F 178 and R242 (Sigma-Genosys), was added to the other tube of PCR master mix to a final primer concentration of 300nM. 45 μ l of the β-Actin or β-GUS master mix was dispensed into wells, in a 96-well plate (Applied BioSystems). 5μ l of the template dilution series was dispensed into triplicate wells for each primer. The reaction was run on an ABI 7900 Sequence Detection System (Applied BioSystems) with the following conditions: 10 min. at 95°C; 40 cycles of

[0438] The Sequence Detection System v2.0 software was used to analyze the fluorescent signal from each well. The high end of the baseline was adjusted to between 8 and 20 cycles to reduce the impact on any data curves, yet be as high as possible to reduce baseline drift. A threshold value was selected that allowed the majority of the amplification curves to cross the threshold during the linear phase of amplification. The disassociation curve for each well was compared to other wells for that marker. This comparison allowed identification of "bad" wells, those that did not amplify, that amplified the wrong size product, or that amplified multiple products. The cycle number at which each amplification curve crossed the threshold (C_T) was recorded and the file transferred to MS Excel for further analysis. The C_T values for triplicate wells were averaged. The data were plotted as a function of the \log_{10} of the calculated starting concentration of RNA. The starting RNA concentration for each cDNA dilution was determined based on the original amount of RNA used in the RT reaction, the dilution of the RT reaction, and the amount used (5 μ l) in the real-time PCR reaction. For each gene, a linear regression line was plotted through all of the dilutions series points. The slope of the line was used to calculate the efficiency of the reaction for each primer set using the equation:

95°C for 15 sec, 60°C for 1 min; followed by a disassociation curve starting at 50°C and ending at 95°C.

$$E = 10^{\left(-\frac{1}{slope}\right)} - 1$$

[0439] Using this equation (Pfaffl 2001, Applied Biosystems User Bulletin #2), the efficiency for these β -actin primers is 1.28 and the efficiency for these β -GUS primers is 1.14 (Figure 10). This efficiency was used when comparing the expression levels among multiple genes and multiple samples. This same method was used to calculate reaction efficiency for primer pairs for each gene studied. A primer pair was considered successful if the efficiency was reproducibly determined to be between 0.7 and 2.4.

SYBR-GREEN ASSAYS

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[0440] Once markers passed the Primer Efficiency QPCR (as stated above), they were used in real-time PCR assays. Patient RNA samples were reverse-transcribed to cDNA (as described above) and 1:10 dilutions made in water. In addition to the patient samples, a no template control (NTC) and a pooled reference RNA (see example 3) described in were included on every plate.

[0441] The Sybr Green real-time PCR reaction was performed using the Taqman Core PCR Reagent kit (Applied BioSystems, Foster City, CA, N808-0228). A master mix was made that consisted of all reagents except the primers and template. The final concentration of all ingredients in the reaction was 1X Taqman Buffer A (Applied BioSystems), 2mM MgCl2 (Applied BioSystems), 200 μ M dATP (Applied BioSystems), 200 μ M dCTP (Applied BioSystems), 200 μ M dGTP (Applied BioSystems), 400 μ M dUTP (Applied BioSystems), 1:400,000 diluted Sybr Green dye (Molecular Probes), 1.25U AmpliTaq Gold (Applied BioSystems). The PCR master mix was aliquotted into eight light-tight tubes, one for each marker to be examined across a set of samples. The optimized primer pair for each marker was then added to the PCR master mix to a final primer concentration of 300nM. 18 μ l of the each marker master mix was dispensed into wells in a 384well plate (Applied BioSystems). 2 μ l of the 1:10 diluted control or patient cDNA sample was dispensed into triplicate wells for each primer pair. The reaction was run on an ABI 7900 Sequence Detection System (Applied BioSystems) using the cycling conditions described above.

[0442] The Sequence Detection System v2.0 software (Applied BioSystems) was used to analyze the fluorescent signal from each well. The high end of the baseline was adjusted to between 8 and 20 cycles to reduce the impact on any data curves, yet be as high as possible to reduce baseline drift. A threshold value was selected that allowed the majority of the amplification curves to cross the threshold during the linear phase of amplification. The disassociation

curve for each well was compared to other wells for that marker. This comparison allowed identification of "bad" wells, those that did not amplify, that amplified the wrong size product, or that amplified multiple products. The cycle number at which each amplification curve crossed the threshold (C_T) was recorded and the file transferred to MS Excel for further analysis. The C_T value representing any well identified as bad by analysis of disassociation curves was deleted. The C_T values for triplicate wells were averaged. A standard deviation (Stdev) and a coefficient of variation (CV) were calculated for the triplicate wells. If the CV was greater than 2, an outlier among the three wells was identified and deleted. Then the average was re-calculated. In each plate, ΔC_T was calculated for each marker-control combination by subtracting the average C_T of the target marker from the average C_T of the control (β -Actin or β -GUS). The expression relative to the control marker was calculated by taking two to the power of the ΔC_T of the target marker. For example, expression relative to β -Actin was calculated by the equation:

$$ErA = 2^{(C_{T,Actin} - C_{T,I \text{ seg es}})}$$

[0443] All plates were run in duplicate and analyzed in the same manner. The percent variation was determined for each sample-marker combination (relative expression) by taking the absolute value of the value of the RE for the second plate from the RE for the first plate, and dividing that by the average. If more than 25% of the variation calculations on a plate are greater than 50%, then a third plate was run.

TAOMAN PROTOCOL

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[0444] Real-time PCR assays were also done using Taqman PCR chemistry.

[0445] The Taqman real-time PCR reaction was performed using the Taqman Universal PCR Master Mix (Applied BioSystems, Foster City, CA, #4324018). The master mix was aliquoted into eight, light-tight tubes, one for each marker. The optimized primer pair for each marker was then added to the correctly labeled tube of PCR master mix. A FAM/ TAMRA dual-labeled Taqman probe (Biosearch Technologies, Navoto, CA, DLO-FT-2) was then added to the correctly labeled tube of PCR master mix. Alternatively, different combinations of fluorescent reporter dyes and quenchers can be used such that the absorption wavelength for the quencher matches the emission wavelength for the reporter, as shown in Table 5. 18µl of the each marker master mix was dispensed into a 384well plate (Applied BioSystems). 2µl of the template sample was dispensed into triplicate wells for each primer pair. The final concentration of each reagent was: 1X TaqMan Universal PCR Master Mix, 300nM each primer, 0.25nM probe, 2µl 1:10 diluted template. The reaction was run on an ABI 7900 Sequence Detection System (Applied Biosystems) using standard conditions (95°C for 10 min., 40 cycles of 95°C for 15 sec, 60°C for 1min.).

[0446] The Sequence Detector v2.0 software (Applied BioSystems) was used to analyze the fluorescent signal from each well. The high end of the baseline was adjusted to between 8 and 20 cycles to reduce the impact on any data curves, yet be as high as possible to reduce baseline drift. A threshold value was selected that allowed most of the amplification curves to cross the threshold during the linear phase of amplification. The cycle number at which each amplification curve crossed the threshold (C_T) was recorded and the file transferred to MS Excel for further analysis. The C_T values for triplicate wells were averaged. The C_T values for triplicate wells were averaged. A standard deviation (Stdev) and a coefficient of variation (CV) were calculated for the triplicate wells. If the CV was greater than 2, an outlier among the three wells was identified and deleted. Then the average was re-calculated. In each plate, ΔC_T was calculated for each marker-control combination by subtracting the average C_T of the target marker from the average C_T of the control (β -Actin, or β -GUS). The expression relative to the control marker was calculated by taking two to the power of the ΔC_T of the target marker. All plates were run in duplicate and analyzed in the same manner. The percent variation was determined for each sample-marker combination (relative expression) by taking the absolute value of the value of the RE for the second plate from the RE for the first plate, and dividing that by the average. If more than 25% of the variation calculations on a plate are greater than 50%, then a third plate was run.

BI-PLEXING

[0447] Variation of real-time PCR assays can arise from unequal amounts of RNA starting material between reactions. In some assays, to reduce variation, the control gene amplification was included in the same reaction well as the target gene. To differentiate the signal from the two genes, different fluorescent dyes were used for the control gene. β -Actin was used as the control gene and the TaqMan probe used was labeled with the fluorescent dye VIC and the quencher TAMRA (Biosearch Technologies, Navoto, CA, DLO-FT-2). Alternatively, other combinations of fluorescent reporter dyes and quenchers (Table 5) can be used as long as the emission wavelength of the reporter for the control gene is sufficiently different from the wavelength of the reporter dye used for the target. The control gene primers and probe

were used at limiting concentrations in the reaction (150 nM primers and 0.125 nM probe) to ensure that there were enough reagents to amplify the target marker. The plates were run under the same protocol and the data are analyzed in the same way, but with a separate baseline and threshold for the VIC signal. Outliers were removed as above from both the FAM and VIC signal channels. The expression relative to control was calculated as above, using the VIC signal from the control gene.

ABSOLUTE QUANTITATION

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[0448] Instead of calculating the expression relative to a reference marker, an absolute quantitation can be performed using real-time PCR. To determine the absolute quantity of each marker, a standard curve is constructed using serial dilutions from a known amount of template for each marker on the plate. The standard curve may be made using cloned genes purified from bacteria or using synthetic complimentary oligonucleotides. In either case, a dilution series that covers the expected range of expression is used as template in a series of wells in the plate. From the average C_T values for these known amounts of template a standard curve can be plotted. From this curve the C_T values for the unknowns are used to identify the starting concentration of cDNA. These absolute quantities can be compared between disease classes (i.e. rejection vs. no-rejection) or can be taken as expression relative to a control gene to correct for variation among samples in sample collection, RNA purification and quantification, cDNA synthesis, and the PCR amplification.

CELL TYPE SPECIFIC EXPRESSION

[0449] Some markers are expressed only in specific types of cells. These markers may be useful markers for differentiation of rejection samples from no-rejection samples or may be used to identify differential expression of other markers in a single cell type. A specific marker for cytotoxic T-lymphocytes (such as CD8) can be used to identify differences in cell proportions in the sample. Other markers that are known to be expressed in this cell type can be compared to the level of CD8 to indicate differential gene expression within CD8 T-cells.

Control genes for PCR

[0450] As discussed above, PCR expression measurements can be made as either absolute quantification of gene expression using a standard curve or relative expression of a gene of interest compared to a control gene. In the latter case, the gene of interest and the control gene are measured in the same sample. This can be done in separate reactions or in the same reaction (biplex format, see above). In either case, the final measurement for expression of a gene is expressed as a ratio of gene expression to control gene expression. It is important for a control gene to be constitutively expressed in the target tissue of interest and have minimal variation in expression on a per cell basis between individuals or between samples derived from an individual. If the gene has this type of expression behavior, the relative expression ratio will help correct for variability in the amount of sample RNA used in an assay. In addition, an ideal control gene has a high level of expression in the sample of interest compared to the genes being assayed. This is important if the gene of interest and control gene are used in a biplex format. The assay is set up so that the control gene reaches its threshold Ct value early and its amplification is limited by primers so that it does not compete for limiting reagents with the gene of interest.

[0451] To identify an ideal control gene for an assay, a number of genes were tested for variability between samples and expression in both mononuclear RNA samples and whole blood RNA samples using the RNA procurement and preparation methods and real-time PCR assays described above. 6 whole-blood and 6 mononuclear RNA samples from transplant recipients were tested. The intensity levels and variability of each gene in duplicate experiments on both sample types are shown in Figure 11.

[0452] Based on criteria of low variability and high expression across samples, β -actin, 18s, GAPDH, b2microglobulin were found to be good examples of control genes for the PAX samples. A single control gene may be incorporated as an internal biplex control is assays.

Controlling for variation in real time PCR

[0453] Due to differences in reagents, experimenters, and preparation methods, and the variability of pipetting steps, there is significant plate-to-plate variation in real-time PCR experiments. This variation can be reduced by automation (to reduce variability and error), reagent lot quality control, and optimal data handling. However, the results on replicate plates are still likely to be different since they are run in the machine at different times.

[0454] Variation can also enter in data extraction and analysis. Real-time PCR results are measured as the time (measured in PCR cycles) at which the fluorescence intensity (\square Rn in Applied Biosystems SDS v2.1 software) crosses a user-determined threshold (CT). When performing relative quantification, the CT value for the target gene is subtracted

from the CT value for a control gene. This difference, called ACT, is the value compared among experiments to determine whether there is a difference between samples. Variation in setting the threshold can introduce additional error. This is especially true in the duplexed experimental format, where both the target gene and the control gene are measured in the same reaction tube. Duplexing is performed using dyes specific to each of the two genes. Since two different fluorescent dyes are used on the plate, two different thresholds are set. Both of these thresholds contribute to each ACT. Slight differences in the each dye's threshold settings (relative to the other dye) from one plate to the next can have significant effects on the ACT.

[0455] There are several methods for setting the threshold for a PCR plate. Older versions of SDS software (Applied Biosystems) determine the average baseline fluorescence for the plate and the standard deviation of the baseline. The threshold is set to 10x the standard deviation of the baseline. In SDS 2.0 the users must set the baseline by themselves. Software from other machine manufacturers either requires the user to set the threshold themselves or uses different algorithms. The latest version of the SDS software (SDS 2.1) contains Automatic baseline and threshold setting. The software sets the baseline separately for each well on the plate using the Δ Rn at cycles preceding detectable levels. Variability among plates is dependent on reproducible threshold setting. This requires a mathematical or experimental data driven threshold setting protocol. Reproducibly setting the threshold according to a standard formula will minimize variation that might be introduced in the threshold setting process. Additionally, there may be experimental variation among plates that can be reduced by setting the threshold to a component of the data. We have developed a system that uses a set of reactions on each plate that are called the threshold calibrator (TCb). The TCb wells are used to set the threshold on all plates.

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- 1. The TCb wells contain a template, primers, and probes that are common among all plates within an experiment.
- 2. The threshold is set within the minimum threshold and maximum threshold determined above.
- 3. The threshold is set to a value in this range that results in the average CT value for the TCb wells to be the same on all plates.

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[0456] These methods were used to derive the primers depicted in Table 2C.

Example 13: Real-time PCR expression markers of acute allograft rejection

[0457] In examples 14 and 16, genes were identified as useful markers of cardiac and renal allograft rejection using microarrays. Some genes identified through these studies arc listed in Table 2. In order to validate these findings, obtain a more precise measurement of expression levels and develop PCR reagents for diagnostic testing, real-time PCR assays were performed on samples from allograft recipients using primers to the identified genes. Some gene specific PCR primers were developed and tested for all genes in Table 2A as described in example 12. Some primers are listed in Table 2C and the sequence listing. These primers were used to measure expression of the genes relative to β -actin or β -gus in 69 mononuclear RNA samples obtained from cardiac allograft recipients using Sybr green real-time PCR assays as described in example 12. Each sample was associated with an ISHLT cardiac rejection biopsy grade. The samples were tested in 2 phases. In phase I, 14 Grade 0, 1 Grade 1A, 3 Grade 2 and 9 Grade 3A samples were tested. In phase II, 19 Grade 2, 4 Grade 1B, 4 Grade 2 and 15 Grade 3A samples were tested. Data was analyzed for each phase individually and for the combined phase I + II sample set. These data are summarized in Table 6.

[0458] The average fold change in expression between rejection (3A) and no rejection (0) samples was calculated. A t-test was done to determine the significance with which each gene was differentially expressed between rejection and no rejection and a p-value was calculated. Genes with high average fold changes and low p-values are considered best candidates for further development as rejection markers. However, it is important to note that a gene with a low average fold change and a high p-value may still be a useful marker for rejection in some patients and may work as part of a gene expression panel to diagnose rejection. These same PCR data were used to create PCR gene expression panels for diagnosis of acute rejection as discussed in example 17.

[0459] Non-parametric tests such as the Fisher Exact Test and Mann-Whitney U test are useful for choosing useful markers. They assess the ability of markers to discrininate between different classes as well as their significance. For example, one could use the median of all samples (including both non-rejector and rejector samples) as a threshold and apply the Fisher Exact test to the numbers of rejectors and non-rejectors above and below the threshold.

[0460] These methods were used to generate the data in Table 2D.

Example 14: Identification of diagnostic nucleotide sets for diagnosis of Cardiac Allograft Rejection using microarrays

[0461] Genes were identified which have expression patterns useful for the diagnosis and monitoring of acute cardiac allograft rejection. Further, sets of genes that work together in a diagnostic algorithm for allograft rejection were identified. Acute allograft rejection is a process that occurs in all solid organ transplantation including, heart, lung, liver, kidney,

pancreas, pancreatic islet cell, intestine and others. Gene expression markers of acute cardiac rejection may be useful for diagnosis and monitoring of all allograft recipients. Patients, patient clinical data and patient samples used in the discovery of markers below were derived from a clinical study described in example 5.

[0462] The collected clinical data was used to define patient or sample groups for correlation of expression data. Patient groups were identified for comparison. For example, a patient group that possesses a useful or interesting clinical distinction, verses a patient group that does not possess the distinction. Measures of cardiac allograft rejection were derived from the clinical data to divide patients (and patient samples) into groups with higher and lower rejection activity over some period of time or at any one point in time. Such data were rejection grades as determined from histological reading of the cardiac biopsy specimens by a pathologist and data measuring progression of end-organ damage, including depressed left ventricular dysfunction (decreased cardiac output, decreased ejection fraction, clinical signs of low cardiac output) and usage of inotropic agents (Kobashigawa 1998). Mononuclear RNA samples were collected and prepared from patients who had recently undergone a cardiac allograft transplantation using the protocol described in example 2. The allograft rejection status at the time of sample collection was determined by examination of cardiac biopsies as described in example 5 and as summarized here.

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[0463] 300 patient samples were included in the analysis. Each patient sample was associated with a biopsy and other clinical data collected at the time of the sample. The cardiac biopsies were graded by a pathologist at the local center and by three centralized pathologists who read the biopsy slides from all four local centers in a blinded manner. Biopsy grades included 0, 1A, 1B, 2, 3A, and 3B. No grade 4 rejection was identified. Dependent variables were developed based on these grades using the local center pathology reading, the reading of a centralized and blinded pathologist, the highest of the readings, local or centralized and a consensus grade derived from all pathological readings. Samples were classified as no rejection or rejection in the following ways: Grade 0 vs. Grades 1-4, Grades 0 and 1A vs. Grades 1B-4, Grade 0 vs. Grade 3A, Grade 0 vs. Grades 1B-4, and Grade 0 vs. Grades 1B and 3A-4. Grade 0 samples were selected such that they were not immediately followed by an episode of acute rejection in the same patient. Comparing Grade 0 samples to Grade 3A samples gives the greatest difference between the rejection and no rejection groups on average.

[0464] Taking the highest of all pathologist readings has the effect of removing any sample from the no rejection class that was not a unanimous Grade 0. It also results in an increase in the number of rejection samples used in an analysis with the assumption that if a pathologist saw features of rejection, the call was likely correct and the other pathologists may have missed the finding. Many leading cardiac pathologists and clinicians believe that ISHLT grade 2 rejection does not represent significant acute rejection. Thus, for correlation analysis, exclusion of Grade 2 samples may be warranted. Clinical data were also used to determine criteria for including samples in the analysis. For example, a patient with an active infection or in the early post-transplant period (ongoing surgical inflammation) might have immune activation unrelated to rejection and thus be difficult to identify as patients without rejection. The strictest inclusion criteria required that samples be from patients who did not have a bacterial or viral infection, were at least two weeks post cardiac transplant, were asymptomatic and were not currently admitted to the hospital.

[0465] After preparation of RNA (example 2), amplification, labeling, hybridization, scanning, feature extraction and data processing were done as described in Example 11, using the oligonucleotide microarrays described in Example 9. The resulting log ratio of expression of Cy3 (patient sample)/ Cy5 (R50 reference RNA) was used for analysis.

[0466] Significance analysis for microarrays (SAM, Tusher 2001, Example 15) was used to discover genes that were differentially expressed between the rejection and no-rejection groups. Many different combinations of dependent variables, inclusion criteria, static/referenced, and data subsets were used in SAM analysis to develop the primary lists of genes significantly differentially expressed between rejection and no-rejection. As described in example 15, SAM assigns a false detection rate to each gene identified as differentially expressed. The most significant of these genes were identified.

[0467] An exemplary analysis was the comparison of Grade 0 samples to Grade 3A-4 samples using SAM. Data from the all the pathological readings was used to identify consensus Grade 0 samples and samples with at least one reading of Grade 3A or above. Using this definition of rejection and no rejection, expression profiles from rejection samples were compared to no rejection samples using SAM. The analysis identified 7 genes with a FDR of 1%, 15 genes @ 1.4%, 35 genes @ 3.9%. Many more genes were identified at higher FDR levels.

[0468] In Table 7, a number of SAM analyses are summarized. In each case the highest grade from the 3 pathologists was taken for analysis. No rejection and rejection classes are defined. Samples are either used regardless of redundancy with respect to patients or a requirement is made that only one sample is used per patient or per patient per class. The number of samples used in the analysis is given and the lowest FDR achieved is noted.

[0469] Some of the genes identified by SAM as candidate rejection markers are noted in Table 2A and B. SAM chooses genes as significantly different based on the magnitude of the difference between the groups and the variation among the samples within each group. It is important to note that a gene which is not identified by SAM as differentially expressed between rejection and no rejection may still be a useful rejection marker because: 1. The microarray technology is not adequately sensitive to detect all genes expressed at low levels. 2. A gene might be a useful member of a gene expression

panel in that it is a useful rejection marker only in a subset of patients. This gene may not be significantly differentially expressed between all rejection and no rejection samples.

[0470] For the purposes of cross-validation of the results, the datasets were also divided into subsets to compare analysis between two subsets of roughly half of the data. The types of subsets constructed were as follows. First half/second half subsets were the first half of the samples and the second half of the samples from a dataset ordered by sample number. Odd/even subsets used the same source, a dataset ordered by sample number, but the odd subset consisted of every 2nd sample starting with the first and the even subset consisted of every 2nd sample starting with the second sample, Center 14/other subsets were the same datasets, divided by transplant hospital. The center 14 subset consisted of all samples from patients at center 14, while the other subset consisted of all samples from the other three centers (12,13, and 15). When a gene was found to be significantly differentially expressed in both sets of data, a higher priority was put on that gene for development of a diagnostic test. This was reflected in a "Array Score" value (Table 2B) that also considered the false detection rate for the gene and the importance of the gene in classification models (see example 17).

[0471] Alternatively one can divide samples into 10 equal parts and do 10-fold cross validation of the results of SAM. [0472] Microarray data was also used to generate classification models for diagnosis of rejection as described in example 17. Genes identified through classification models as useful in the diagnosis of rejection are noted in Table 2B in the column "models".

[0473] As genes were identified as useful rejection markers by microarray significance analysis, classification models, PCR analysis, or through searching the prior art, a variety of approaches were employed to discover genes that had similar expression behavior (coexpression) to the gene of interest. If a gene is a useful rejection marker, then a gene that is identified as having similar expression behavior is also likely to be a useful rejection marker. Hierarchical clustering (Eisen et al. 1998, see example 15) was used to identify co-expressed genes for established rejection markers. Genes were identified from the nearest branches of the clustering dendrogram. Gene expression profiles generated from 240 samples derived from transplant recipients were generated as described above. Hierarchical clustering was performed and co-expressed genes of rejection markers were identified. An example is shown in Figure 12. SEQ ID NO:85 was shown to be significantly differentially expressed between rejection and no rejection using both microarrays and PCR. Gene SEQ ID NO:3020 was identified by hierarchical clustering as closely co-expressed with SEQ ID NO:85.

[0474] Some of the primers for real-time PCR validation were designed for each of the marker genes as described in Example 12 and are listed in Table 2C and the sequence listing. PCR expression measurements using these primers were used to validate array findings, more accurately measure differential gene expression and create PCR gene expression panels for diagnosis of rejection as described in example 17.

[0475] Alternative methods of analyzing the data may involve 1) using the sample channel without normalization by the reference channel, 2) using an intensity-dependent normalization based on the reference which provides a greater correction when the signal in the reference channel is large, 3) using the data without background subtraction or subtracting an empirically derived function of the background intensity rather than the background itself.

[0476] These methods were used to identify genes listed in Table 2B.

Example 15: Correlation and Classification Analysis

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[0477] After generation and processing of expression data sets from microarrays as described in Example 11, a log ratio value is used for most subsequent analysis. This is the logarithm of the expression ratio for each gene between sample and universal reference. The processing algorithm assigns a number of flags to data that are of low signal to noise, saturated signal or are in some other way of low or uncertain quality. Correlation analysis can proceed with all the data (including the flagged data) or can be done on filtered data sets where the flagged data is removed from the set. Filtered data should have less variability and noise and may result in more significant or predictive results. Flagged data contains all information available and may allow discovery of genes that are missed with the filtered data set. After filtering the data for quality as described above and in example 11, missing data are common in microarray data sets. Some algorithms don't require complete data sets and can thus tolerate missing values. Other algorithms are optimal with or require imputed values for missing data. Analysis of data sets with missing values can proceed by filtering all genes from the analysis that have more than 5%, 10%, 20%, 40%, 50%, 60% or other % of values missing across all samples in the analysis. Imputation of data for missing values can be done by a variety of methods such as using the row mean, the column mean, the nearest neighbor or some other calculated number. Except when noted, default settings for filtering and imputation were used to prepare the data for all analytical software packages.

[0478] In addition to expression data, clinical data are included in the analysis. Continuous variables, such as the ejection fraction of the heart measured by echocardiography or the white blood cell count can be used for correlation analysis. Any piece of clinical data collected on study subjects can be used in a correlation or classification analysis. In some cases, it may be desirable to take the logarithm of the values before analysis. These variables can be included in an analysis along with gene expression values, in which case they are treated as another "gene". Sets of markers can

be discovered that work to diagnose a patient condition and these can include both genes and clinical parameters. Categorical variables such as male or female can also be used as variables for correlation analysis. For example, the sex of a patient may be an important splitter for a classification tree.

[0479] Clinical data are used as supervising vectors (dependent variables) for the significance or classification analysis of expression data. In this case, clinical data associated with the samples are used to divide samples in to clinically meaningful diagnostic categories for correlation or classification analysis. For example, pathologic specimens from kidney biopsies can be used to divide lupus patients into groups with and without kidney disease. A third or more categories can also be included (for example "unknown" or "not reported"). After generation of expression data and definition of supervising vectors, correlation, significance and classification analysis are used to determine which set of genes and set of genes are most appropriate for diagnosis and classification of patients and patient samples. Two main types of expression data analyses are commonly performed on the expression data with differing results and purposes. The first is significance analyses or analyses of difference. In this case, the goal of the analysis is to identify genes that are differentially expressed between sample groups and to assign a statistical confidence to those genes that are identified. These genes may be markers of the disease process in question and are further studied and developed as diagnostic tools for the indication. The second major type of analysis is classification analysis. While significance analysis identifies individual genes that are differentially expressed between sample groups, classification analysis identifies gene sets and an algorithm for their gene expression values that best distinguish sample (patient) groups. The resulting gene expression panel and algorithm can be used to create and implement a diagnostic test. The set of genes and the algorithm for their use as a diagnostic tool are often referred to herein as a "model". Individual markers can also be used to create a gene expression diagnostic model. However, multiple genes (or gene sets) are often more useful and accurate diagnostic tools.

Significance analysis for microarrays (SAM)

[0480] Significance analysis for microarrays (SAM) (Tusher 2001) is a method through which genes with a correlation between their expression values and the response vector are statistically discovered and assigned a statistical significance. The ratio of false significant to significant genes is the False Discovery Rate (FDR). This means that for each threshold there are some number of genes that are called significant, and the FDR gives a confidence level for this claim. If a gene is called differentially expressed between two classes by SAM, with a FDR of 5%, there is a 95% chance that the gene is actually differentially expressed between the classes. SAM will identify genes that are differentially expressed between the classes. The algorithm selects genes with low variance within a class and large variance between classes. The algorithm may not identify genes that are useful in classification, but are not differentially expressed in many of the samples. For example, a gene that is a useful marker for disease in women and not men, may not be a highly significant marker in a SAM analysis, but may be useful as part of a gene set for diagnosis of a multi-gene algorithm.

[0481] After generation of data from patient samples and definition of categories using clinical data as supervising vectors, SAM is used to detect genes that are likely to be differentially expressed between the groupings. Those genes with the highest significance can be validated by real-time PCR (Example 13) or can be used to build a classification algorithm as described here.

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[0482] Classification algorithms are used to identify sets of genes and formulas for the expression levels of those genes that can be applied as diagnostic and disease monitoring tests. The same classification algorithms can be applied to all types of expression and proteomic data, including microarray and PCR based expression data. Examples of classification models are given in example 17. The discussion below describes the algorithms that were used and how they were used.

[0483] Classification and Regression Trees (CART) is a decision tree classification algorithm (Breiman 1984). From gene expression and or other data, CART can develop a decision tree for the classification of samples. Each node on the decision tree involves a query about the expression level of one or more genes or variables. Samples that are above the threshold go down one branch of the decision tree and samples that are not go down the other branch. Genes from expression data sets can be selected for classification building with CART by significant differential expression in SAM analysis (or other significance test), identification by supervised tree-harvesting analysis, high fold change between sample groups, or known relevance to classification of the target diseases. In addition, clinical data can be used as independent variables for CART that are of known importance to the clinical question or are found to be significant predictors by multivariate analysis or some other technique. CART identifies predictive variables and their associated decision rules for classification (diagnosis). CART also identifies surrogates for each splitter (genes that are the next best substitute for a useful gene in classification). Analysis is performed in CART by weighting misclassification costs to optimize desired performance of the assay. For example, it may be most important that the sensitivity of a test for a

given diagnosis be > 90%. CART models can be built and tested using 10 fold cross-validation or v-fold cross validation (see below). CART works best with a smaller number of variables (5-50). Multiple Additive Regression Trees (Friedman, JH 1999, MART) is similar to CART in that it is a classification algorithm that builds decision trees to distinguish groups. MART builds numerous trees for any classification problem and the resulting model involves a combination of the multiple trees. MART can select variables as it build models and thus can be used on large data sets, such as those derived from an 8000 gene microarray. Because MART uses a combination of many trees and does not take too much information from any one tree, it resists over training. MART identifies a set of genes and an algorithm for their use as a classifier. [0484] A Nearest Shrunken Centroids Classifier can be applied to microarray or other data sets by the methods described by Tibshirani et al. 2002. This algorithms also identified gene sets for classification and determines their 10 fold cross validation error rates for each class of samples. The algorithm determines the error rates for models of any size, from one gene to all genes in the set. The error rates for either or both sample classes can are minimized when a particular number of genes are used. When this gene number is determined, the algorithm associated with the selected genes can be identified and employed as a classifier on prospective sample.

[0485] For each classification algorithm and for significance analysis, gene sets and diagnostic algorithms that are built are tested by cross validation and prospective validation. Validation of the algorithm by these means yields an estimate of the predictive value of the algorithm on the target population. There are many approaches, including a 10 fold cross validation analysis in which 10% of the training samples are left out of the analysis and the classification algorithm is built with the remaining 90%. The 10% are then used as a test set for the algorithm. The process is repeated 10 times with 10% of the samples being left out as a test set each time. Through this analysis, one can derive a cross validation error which helps estimate the robustness of the algorithm for use on prospective (test) samples. Any % of the samples can be left out for cross validation (v-fold cross validation, LOOCV). When a gene set is established for a diagnosis with an acceptable cross validation error, this set of genes is tested using samples that were not included in the initial analysis (test samples). These samples may be taken from archives generated during the clinical study. Alternatively, a new prospective clinical study can be initiated, where samples are obtained and the gene set is used to predict patient diagnoses.

Example 16: Acute allograft rejection: biopsy tissue gene expression profiling

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[0486] Acute allograft rejection involves activation of recipient leukocytes and infiltration into the rejecting organ. For example, CD8 T-cells are activated by CD4 T-cells and enter the allograft where they destroy graft tissue. These activated, graft-associated leukocytes may reside in the graft, die or exit the graft. Upon exiting, the cells can find their way into the urine or blood (in the case of renal allografts), bile or blood (liver allografts) or blood (cardiac allografts). These activated cells have specific gene expression patterns that can be measured using microarrays, PCR or other methods. These gene expression patterns can be measured in the graft tissue (graft associated leukocytes), blood leukocytes, urine leukocytes or stool/biliary leukocytes. Thus graft associated leukocyte gene expression patterns are used to discover markers of activated leukocytes that can be measured outside the graft for diagnostic testing.

[0487] Renal biopsy and cardiac biopsy tissue specimens were obtained for gene expression profiling. The specimens were obtained at the time of allograft biopsy and were preserved by flash freezing in liquid nitrogen using standard approaches or immersion in an RNA stablization reagent as per the manufacturers recommendation (RNAlater, Qiagen, Valencia, CA). Biopsy allograft pathological evaluation was also obtained and samples were classified as having a particular ISHLT rejection grade (for cardiac) or acute rejection, chronic rejection, acute tubular necrosis or no disease (for renal)

[0488] 28 renal biopsy tissue samples were transferred to RLT buffer, homogenized and RNA was prepared using RNeasy preparation kits (Qiagen, Valencia, CA). Average total RNA yield was 1.3 ug. Samples were subjected to on column DNAse digestion. 18 samples were derived from patients with ongoing acute allograft rejection and 10 were from controls with chronic rejection or acute renal failure.

[0489] RNA from the samples was used for amplification, labeling and hybridization to leukocyte arrays (example 11). Significance analysis for microarrays (SAM, Tusher 2001, Example 15) was used to identify genes that were differentially expressed between the acute rejection samples and controls. Leukocyte markers of acute rejection that are associated with the graft should be genes that are expressed at some level in activated leukocytes. Since leukocytes appear in graft tissue with some frequency with acute rejection, leukocyte genes associate with rejection are identified by SAM as upregulated in acute rejection in this experiment. 35 genes were identified as upregulated in acute rejection by SAM with less than a 5% false detection rate and 139 were detected with < 10.0% FDR.

[0490] For each of these genes, to 50mer oligonucleotide sequence was used to search NCBI databases including Unigene and OMIM. Genes were identified by sequence analysis to be either known leukocyte specific markers, known leukocyte expressed markers, known not to be leukocyte expressed or expression unknown. This information helped selected candidate leukocyte markers from all upregulated genes. This is necessary because some of the upregulated genes may have been expressed by renal tissue. Those genes that are leukocyte specific or leukocyte expressed were

selected for evaluation by PCR in urine and blood samples from patients with and without acute allograft rejection (cardiac and renal). These genes are useful expression markers of acute rejection in allograft tissue specimens and may also be useful gene expression markers for the process in circulating leukocytes, or urine leukocytes. In addition, some of the leukocyte expressed genes from this analysis were selected for PCR validation and development for diagnosis of acute cardiac rejection and are noted in Table 2.

[0491] Five cardiac rejection markers in the peripheral blood were assayed using real-time PCR in renal biopsy specimens. The average fold change for these genes between acute rejection (n = 6) and controls (n = 6) is given below. Work is ongoing to increase the number of samples tested and the significance of the results.

[0492] PCR assays of cardiac rejection peripheral blood markers in renal allograft tissue. R= rejection, NR = No rejection.

Gene	Fold change (R/NR)
Granzyme B	2.16
CD20	1.42
NK cell receptor	1.72
T-box 21	1.74
IL4	1.3

[0493] Markers of renal rejection that are secreted from cells may be measured in the urine or serum of patients as a diagnostic or screening assay for rejection. Genes with lower molecular weight are most likely to be filtered into the urine to be measured in this way. Standard immunoassays may be used to measure these proteins.

Example 17: Microarray and PCR gene expression panels for diagnosis and monitoring of acute allograft rejection

Array panels / classification models

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[0494] Using the methods of the invention, gene expression panels were discovered for screening and diagnosis of acute allograft rejection. Gene expression panels can be implemented for diagnostic testing using any one of a variety of technologies, including, but not limited to, microarrays and real-time PCR.

[0495] Using peripheral blood mononuclear cell RNA that was collected and prepared from cardiac allograft recipients as described in examples 2 and 5, leukocyte gene expression profiles were generated and analyzed using microarrays as described in examples 11, 13, and 15. 300 samples were analyzed. ISHLT rejection grades were used to divide patients into classes of rejection and no rejection. Multiple Additive Regression Trees (MART, Friedman, JH 1999, example 15) was used to build a gene expression panel and algorithm for the diagnosis of rejection with high sensitivity. Default settings for the implementaion of MART called TreeNet 1.0 (Salford Systems, San Diego, CA) were used except where noted.

[0496] 82 Grade 0 (rejection) samples and 76 Grade 1B-4 (no rejection) samples were divided into training (80% of each class) and testing (20% of each class) sets. A MART algorithm was then developed on the training set to distinguish rejection from no rejection samples using a cost of 1.02:1 for misclassification of rejection as no rejection. The resulting algorithm was then used to classify the test samples. The algorithm correctly classified 51 of 66 (77%) no rejection samples in the training set and 9 of 16 (56%) no rejection samples in the test set. For rejection samples 64 of 64 (100%) were correctly classified in the training set and 12 of 12 were correctly classified in the test set. The algorithm used 37 genes. MART ranks genes by order of importance to the model. In order, the 37 genes were: SEQ IDs: 3058, 3030, 3034, 3069, 3081, 3072, 3041, 3052, 3048, 3045, 3059, 3075, 3024, 279, 3023, 3053, 3022, 3067, 3020, 3047, 3033, 3068, 3060, 3063, 3028, 3032, 3025, 3046, 3065, 3080,3039,3055,49,3080,3038,3071.

[0497] Another MART model was built by excluding samples derived from patients in the first month post transplant and from patients with known CMV infection. 20 Grade 0 (rejection) samples and 25 Grade 1B-4 (no rejection) samples were divided into training (80% of each class) and testing (20% of each class) sets. A MART algorithm was then developed on the training set to distinguish rejection from no rejection samples using default settings. The resulting algorithm was then used to classify the test samples. The algorithm correctly classified 100% of samples of both classes in the training and testing sets. However, this model required 169 genes. The sample analysis was done a second time with the only difference being requirement that all decision trees in the algorithm be composed of two nodes (single decision, "stump model"). In this case 15/16 no rejection samples were correctly identified in the training set and 4/4 no rejection samples were correctly identified in the test set. For the rejection samples, 17/19 were correctly identified in the training set and 5/6 were correctly classified in the test set. This model required 23 genes. In order of importance, they were: SEQ IDs:

3042, 2783, 3076, 3029, 3026, 2751, 3036, 3073, 3035, 3050, 3051, 3027, 3074, 3062, 3044, 3077, 2772,3049,3043,3079,3070,3057,3078.

Real-time PCR panels / classification models

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[0498] PCR primers were developed for top rejection markers and used in real-time PCR assays on transplant patient samples as described in examples 12 and 13. This data was used to build PCR gene expression panels for diagnosis of rejection. Using MART (example 15) a 10-fold cross validated model was created to diagnose rejection using 12 no rejection samples (grade 0) and 10 rejection samples (grade 3A). Default settings were used with the exception of assigning a 1.02:1 cost for misclassification of rejection as no rejection and requirement that all decision trees be limited to 2 nodes ("stump model"). 20 genes were used in the model, including: SEQ IDs:101, 3021, 102, 2781, 78, 87, 86, 36, 77, 2766, 3018, 80, 3019, 2752, 79, 99, 3016, 2790, 3020, 3056, 88. The 10-fold cross-validated sensitivity for rejection was 100% and the specificity was 85%. Some PCR primers for the genes are listed in Table 2C and the sequence listing.

[0499] A different analysis of the PCR data was performed using the nearest shrunken centroids classifier (Tibshirani et al. 2002; PAM version 1.01, see example 15). A 10-fold cross validated model was created to diagnose rejection using 13 no rejection samples (grade 0) and 10 rejection samples (grade 3A). Default settings were used with the exception of using a prior probability setting of (0.5, 0.5). The algorithm derives algorithms using any number of the genes. A 3-gene model was highly accurate with a 10 fold cross-validated sensitivity for rejection of 90%, and a specificity of 85%.

[0500] The 3 genes used in this model were: SEQ IDs 2784, 79, and 2794. Some of the PCR primers used are given in Table 2C and the sequence listing. An ROC curve was plotted for the 3-gene model and is shown in Figure 13.

Example 18: Assay sample preparation

[0501] In order to show that XDx's leukocyte-specific markers can be detected in whole blood, we collected whole blood RNA using the PAXgene whole blood collection, stabilization, and RNA isolation kit (PreAnalytix). Varying amounts of the whole blood RNA were used in the initial RT reaction (1, 2, 4, and 8ug), and varying dilutions of the different RT reactions were tested (1:5, 1:10, 1:20, 1:40, 1:80, 1:160). We did real-time PCR assays with primers specific to XDx's markers and showed that we can reliably detect these markers in whole blood.

[0502] Total RNA was prepared from 14 mononuclear samples (CPT, BD) paired with 14 whole blood samples (PAX-gene, PreAnalytix) from transplant recipients, cDNA was prepared from each sample using 2ug total RNA as starting material. Resulting cDNA was diluted 1:10 and Sybr green real-time PCR assays were performed.

[0503] For real-time PCR assays, Ct values of 15-30 are desired for each gene. If a gene's Ct value is much above 30, the result may be variable and non-linear. For PAX sample, target RNA will be more dilute than in CPT samples, cDNA dilutions must be appropriate to bring Ct values to less than 30. Ct values for the first 5 genes tested in this way are shown in the table below for both whole blood RNA (PAX) and mononuclear RNA (CPT).

Gene	Ct PAX	Ct CPT
CD20	27.41512	26.70474
4761	28.45656	26.52635
3096	29.09821	27.83281
Granzyme B	31.18779	30.56954
IL4	33.11774	34.8002
Actin	19.17622	18.32966
B-GUS	26.89142	26.92735

[0504] With one exception, the genes have higher Ct values in whole blood. Using this protocol, all genes can be detected with Cts <35. For genes found to have Ct values above 30 in target samples, less diluted cDNA may be needed.

Example 19: Allograft rejection diagnostic gene sequence analysis

[0505] Gene products that are secreted from cells or expressed as surface proteins have special diagnostic utility in

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that an assay may be developed to detect relative quantities of proteins in blood plasma or serum. Secreted proteins may also be detectable in urine, which may be a useful sample for the detection of rejection in renal allograft recipients. Cell surface markers may be detected using antigen specific antibodies in ELISA assays or using flow srting techniques such as FACS.

[0506] Each gene that is found to be differentially regulated in one population of patients has several potential applications. It may be a target for new pharmaceuticals, a diagnostic marker for a condition, a benchmark for titrating drug delivery and clearance, or used in screening small molecules for new therapeutics. Any of these applications may be improved by an understanding of the physiologic function and localization of the gene product in vivo and by relating those functions to known diseases and disorders. Identifying the basic function of each candidate gene helps identify the signaling or metabolic pathways the gene is a part of, leading us to investigate other members of those pathways as potential diagnostic markers or targets of interest to drug developers.

[0507] For each of the markers in table 2, we attempted to identify the basic function and subcellular localization of the gene. These results are summarized in Table 9. In addition to initial DNA sequencing and processing, sequence analysis, and analysis of novel clones, information was obtained from the following public resources: Online Mendelian Inheritance in Man at the NCBI, LocusLink at the NCBI, the SWISS-PROT database, and Protein Reviews on the Web. For each marker represented by a curated reference mRNA from the RefSeq project, the corresponding reference protein accession number is listed. Curated sequences are those that have been manually processed by NCBI staff to represent the best estimate of the mRNA sequence as it is transcribed, based on alignments of draft DNA sequence, predicted initiation, termination and splice sites, and submissions of EST and full-length mRNA sequences from the scientific community.

[0508] These methods were used to derive the data in Table 2E.

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Example 20: Detection of proteins expressed by diagnostic gene sequences

[0509] One of ordinary skill in the art is aware of many possible methods of protein detection. The following example illustrates one possible method.

[0510] The designated coding region of the sequence is amplified by PCR with adapter sequences at either end for subcloning. An epitope or other affinity "tag" such as a "His-tag" may be added to facilitate purification and/or detection of the protein. The amplified sequence is inserted into an appropriate expression vector, most typically a shuttle vector which can replicate in either bacteria, most typically E. coli, and the organism/cell of choice for expression such as a yeast or mammalian cell. Such shuttle vectors typically contain origins of replication for bacteria and an antibiotic resistance marker for selection in bacteria, as well as the relevant replication and selection sequences for transformation/ transfection into the ultimate expression cell type. In addition, the sequence of interest is inserted into the vector so that the signals necessary for transcription (a promoter) and translation operably linked to the coding region. Said expression could be accomplished in bacteria, fungi, or mammalian cells, or by in vitro translation.

[0511] The expression vector would then typically be used to transform bacteria and clones analyzed to ensure that the proper sequence had been inserted into the expression vector in the productive orientation for expression. Said verified expression vector is then transfected into a host cell and transformants selected by a variety of methods including antibiotic resistance or nutritional complementation of an auxotrophic marker. Said transformed cells are then grown under conditions conducive to expression of the protein of interest, the cells and conditioned media harvested, and the protein of interest isolated from the most enriched source, either the cell pellet or media.

[0512] The protein is then be isolated by standard of chromatographic or other methods, including immunoaffinity chromatography using the affinity "tag" sequence or other methods, including cell fractionation, ion exchange, size exclusion chromatography, or selective precipitation. The isolated and purified protein is then be used as an antigen to generate specific antibodies. This is accomplished by standard methods including injection into heterologous species with an adjuvant, isolation of monoclonal antibodies from mice, or in vitro selection of antibodies from bacteriophage display antibody libraries. These antibodies are then used to detect the presence of the indicated protein of interest in a complex bodily fluid using standard methods such as ELISA or RIA.

Example 21: Detecting changes in the rate of hematopoiesis

[0513] Gene expression profiling of blood cells from cardiac allograft recipients was done using microarrays and real-time PCR as described in other examples herein.

[0514] Two of the genes in that were most correlated with cardiac transplant acute rejection with both microarrays and PCR were hemoglobin Beta and 2,3 DPGM. These genes are well know to be specific markers of erythrocyte lineages. This correlation was found using both purified peripheral mononuclear cells and whole blood RNA preparations. **[0515]** Analysis of the five genes from the PCR data most strongly correlated with rejection showed that their expression levels were extremely highly correlated within each other (R2 > 0.85).

Gene	Hs	Acc	SEQ ID No
hemoglobin, beta (HBB)	Hs.155376	NM_000518	86
2,3-bisphosphoglycerate mutase (BP	Hs.198365	X04327	87
cDNA FLJ20347	Hs.102669	AK000354	94
1602620663F1cDNA	Hs.34549	AI123826	107
HA 1247 cDNA	Hs.33757	AI114652	91

[0516] This suggested that they were all elevated as part of a single response or process. When the microarray data was used to cluster these genes with each other and the other genes on the microarray, we found that these five genes clustered reasonably near each and of the other array genes which clustered tightly with them, four of the top 40 or so were platelet related genes. In addition, these a number of these genes clustered closely with CD34. CD34 is a marker of hematopoietic stem cells and is seen in the peripheral blood with increased hematopoisis.

[0517] CD34, platelet RNA and erythrocyte RNA all mark immature or progenitor blood cells and it is clear that theses marker of acute rejection are part of a coordinated hematopoietic response. A small increase in the rate of production of RBCs and platelets may result in large fold changes in RNA levels. Immune activation from acute rejection may lead to increased hamatopoiesis in the bone marrow and non-marrow sites. This leads to an increase in many lineages because of the lack of complete specificity of the marrow response. Alternatively, increased hematopoiesis may occur in a transplant recipient due to an infection (viral or other), allergy or other stimulus to the system. This results in production of cells or a critical mass of immune cells that can cause rejection. In this scenario, monitoring for markers of immune activation would provide an opportunity for early diagnosis.

Table 1

Disease Classification	Disease/Patient Group
	•
Cardiovascular Disease	Atherosclerosis
	Unstable angina
	Myocardial Infarction
	Restenosis after angioplasty
	Congestive Heart Failure
	Myocarditis
	Endocarditis
	Endothelial Dysfunction
	Cardiomyopathy
	Cardiovascular drug use
Infectious Disease	Hepatitis A, B, C, D, E, G
	Malaria
	Tuberculosis
	HIV
	Pneumocystis Carinii
	Giardia
	Toxoplasmosis
	Lyme Disease
	Rocky Mountain Spotted Fever
	Cytomegalovirus
	Epstein Barr Virus
	Herpes Simplex Virus
	Clostridium Dificile Colitis
	Meningitis (all organisms)
	Pneumonia (all organisms)
	Urinary Tract Infection (all organisms)
	Infectious Diarrhea (all organisms)

(continued)

	Disease Classification	Disease/Patient Group
		Anti-infectious drug use
5	Angiogenesis	Pathologic angiogenesis Physiologic angiogenesis Treatment induced angiogenesis Pro or anti-angiogenic drug use
10	Transplant Rejection	Heart Lung Liver
15		Pancreas Bowel Bone Marrow Stem Cell Graft versus host disease
20		Transplant vasculopathy Skin Cornea Islet Cells
25		Kidney Xenotransplants Mechanical Organ Immunosupressive drug use
30	Hematological Disorders	Anemia - Iron Deficiency Anemia - B12, Folate deficiency Anemia-Aplastic Anemia - hemolytic Anemia - Renal failure
35		Anemia - Chronic disease Polycythemia rubra vera Pernicious anemia Idiophic Thrrombocytopenic purpura Thrombotic Thrombocytopenic purpura
40		Essential thrombocytosis Leukemia Cytopenias due to immunosupression Cytopenias due to Chemotherapy Myelodysplasia

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10		SEQ ID RNA/cDNA																								
15		SEQ ID																								
20		Acc	X01205	NM 001101	NM_000181	NM 004048	NM 004622	NM_001838	NM_004633	NM_004847	NM_000032.1	NM_017413	NM_005191	NM_004437	NM_004351	625000_MN	NM_000902	NM_002259	695000 ⁻ MN	0V_000570	NM_002286	NM_000442	NM_001773	995000_MN	NM_003234	NM_001836
25																										
30	Table 2A.	HS	NA	Hs.288061	Hs.183868	Hs.75415	Hs.75066	Hs.1652	Hs.25333	Hs.76364	Hs.323383	Hs.303084	Hs.838	Hs.37427	Hs.3144	Hs.54443	Hs.1298	Hs.74082	Hs.176663	Hs.372679	Hs.74011	Hs.78146	Hs.374990	Hs.77424	Hs.77356	Hs.135626
35		SEQ ID 50mer																								
40										tor 1, all															eptor	
45			al RNA		e, beta	ylobulin				mmatory fac															ansferrin rec	
50		Gene Name	18S ribosomal RNA	Actin, beta	Glucuronidase, beta	beta 2 microglobulin	Translin	1707	4685-IL1R	Allograft inflammatory factor 1, all variants	ALAS2	APELIN	B7-1, CD80	Band 4.1	c-cbl-B	CCR5	CD10	CD159a	CD16	CD16b	CD223	CD31	CD34	CD64	CD71 =T9, transferrin receptor	chymase
55		Gene	HSRRN18S	АСТВ	GUSB	B2M	TSN	CCR7	IL1R2	AIF-1	ALAS2	APELIN	CD80	EPB41	CBLB	CCR5	MME	KLRC1	FCGR3A	FCGR3B	LAG3	PECAM1	CD34	FCGR1A	TFRC	CMA1

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10		SEQ ID RNA/cDNA												89											
15		S												368											
20		ACC	NM_000222	NM_005373	NM_004445	NM_000121.2	NM_014009	NM_002049	NM_000419	NM_006433	NM_006144	NM_000558.3	NM_005332.2	NM_000518.4	NM_000519.2	NM_005330	NM_000559.2	NM_005331	NM_033554	NM_002122	NM_002124.1	NM_012092	NM_001562	NM_000588	NM_000885
25																									
30	(continued)	HS	Hs.81665	Hs.84171	Hs.3796	Hs.127826	Hs.247700	Hs.765	NM 000419.2	Hs.105806	Hs.90708	Hs.398636	Hs.272003	Hs.155376	Hs.36977	Hs.117848	Hs.283108	Hs.247921	Hs.198253	Hs.198253	Hs.375570	Hs.56247	Hs.83077	Hs.694	Hs.40034
35		SEQ ID 50mer												36											
40																								ılating	CD49D,
45												alpha 1	zeta	beta	delta	epsilon 1	gamma A	theta 1	II, DP alpha 1	MHC, class II, DQ alpha 1	I, DR beta I			interleukin 3 (colony-stimulating factor, multiple)	Integrin, alpha 4 (antigen CD49D,
50		Gene Name	c-Kit	c-mpl	EphB6	EPO-R	Foxp3	GATA1	all 95	granulysin	GZMA	hemoglobin, alpha 1	hemoglobin, zeta	hemoglobin, beta	hemoglobin, delta	hemoglobin, epsilon 1	hemoglobin, gamma A	hemoglobin, theta 1	MH/c, class II, DP alpha	MHC, class I	MHC, class II, DR beta	SOOI	IL18	interleukin 3 (cc factor, multiple)	Integrin, alph
55		Gene	KIT	MPL	EphB6	EPOR	Foxp3	GATA1	ITGA2B	GNLY	GZMA	HBA	HBZ	HBB	НВD	HBE	HBG	HBQ	HLA-DP	HLA-DQ	HLA-DRB	ICOS	IL18	1.3	ITGA4

55	50	45	40	35	30	25	20	15	10	5
					(continued)					
Gene	Gene Name			SEQ ID 50mer	HS		ACC	SEQ ID R	SEQ ID RNA/cDNA	
ITGAM	integrin, alpha M (complem component receptor 3, alpl knownas CD11b (p170), m: antigen alpha polypeptide)	integrin, alpha M (complement component receptor 3, alpha; also known as CD11b (p170), macrophage antigen alpha polypeptide)	e g		Hs.172631		NM_000632			
ITGB7	integrin, beta 7			49	Hs.1741		000889 WN	381		
CEBPB	LAP, CCAAT/enhance protein (C/EBP), beta	LAP, CCAAT/enhancer binding protein (C/EBP), beta			Hs.99029		NM_005194			
NF-E2	NF-E2				Hs.75643 Hs.158297	8297	NM 006163			
PDCD1	programmed ce	programmed cell death 1, PD-1					NM_005018			
PF4	platelet factor 4 motif) ligand 4)	platelet factor 4 (chemokine (C-X-C motif) ligand 4)	O		Hs.81564		NM_002619			
PRKCQ	protein kinase C, theta	C, theta			Hs.211593 Hs.198468	98468	NM_006257.1			
PPARGC1	PPARgamma						NM 013261			
RAG1	recombination &	recombination activating gene 1			Hs.73958		NM 000448			
RAG2	recombination a	recombination activating gene 2			Na		NM 000536			
CXCL12	chemokine (C-) (stromal cell-de	chemokine (C-X-C motif) ligand 12 (stromal cell-derived factor 1) (SDF-1)	7		Hs.237356		609000 ⁻ WN			
TNFRSF4	tumor necrosis factor receptor superfamily, member 4	factor receptor ember 4			Hs.129780 Hs.181097	81097	NM_003327			
TNFSF4	tumor necrosis factor (ligand) superfamily, member 4 (tax-transcriptionally activated glycoprotein 1, 34kDa)	factor (ligand) ember 4 onally activated 34kDa)					NM_003326			
TIPS1	tryptase, alpha				Hs.334455		NM 003293			
ADA	ADA adenosine deaminase	e deaminase			Hs.1217		NM 000022			
CPM	Carboxypeptidase M	ase M			Hs.334873		NM 001874.1			
CSF2	colony stimulati	colony stimulating factor, GM-CSF			Hs.1349		NM 000758.2			
CSF3	colony stimulati	colony stimulating factor 3, G-CSF			Hs.2233		NM 172219			

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5																				
10		SEQ ID RNA/cDNA																		
15		SEQ												409	410	411	412			
20		Acc	NM_000567.1	NM_004119	NM 002051.1	NM 002185.1	NM_006563.1	NM_005356.2	NM_016269.2	NM_002659.1	NM_006573.3	NM 000584	NM_004131	NM_000639	NM_006019	NM_005041	000589 NM_000589	NM_002188	NM_005214	NM_001768
25																				
30	(continued)	HS	Hs.76452	Hs.385	Hs.169946	Hs.362807	Hs37860	Hs.1765	Hs.44865	Hs.179657	Hs.270737	Hs.624	Hs.1051	Hs.2007	Hs.46465	Hs.2200	Hs.73917	Hs.845	Hs.247824	Hs.85258
35		SEQ ID 50mer												77	78	79	80			
40			axin-related	Kinase 3			Kruppel-like factor I (erythroid), EKLF	tein tyrosine	ling factor 1	ogen 37, uPAR	igand) 8b,		2, cytotoxic d serine	igand)	T-cell, immune regulator 1, ATPase, H+ transporting, lysosomal V0 protein a isoform 3	protein)			-associated	CD8 antigen, alpha polypeptide (p32)
45		Ф	C-reactive protein, pentraxin-related (CRP),	FMS-Related Tyrosine Kinase	GATA binding protein 3	7 receptor	e factor I (ery	lymphocyte-specific protein tyrosine kinase	lymphoid enhancer-binding factor	Urokinase-type Plasminogen Activator Receptor, CD87, uPAR	Tumor necrosis factor (ligand) superfamily, member 13b, BlyS/TALL-1/BAFF	8	Granzyme B (granzyme 2, cytotoxic T-lymphocyte-associated serine esterase 1)	Tumor necrosis factor (ligand) superfamily, member 6	T-cell, immune regulator 1, ATPase, H+ transporting, lysosomal V0 proteir a isoform 3	Perforin 1 (pore forming protein)	4	13	Cytotoxic T-lympbocyte-associated protein 4	n, alpha poly
50		Gene Name	C-reactive p	FMS-Relate	GATA bindi	Interleukin 7 receptor	Kruppel-lik€	lymphocyte kinase	lymphoid ei	Urokinase-i Activator Re	Tumor necrosis fac superfamily, membr BlyS/TALL-1/BAFF	Interleukin 8	Granzyme F T-lymphocy esterase 1)	Tumor neci superfamily	T-cell, immu H+ transpor a isoform 3	Perforin 1 (Interleukin 4	Interleukin 13	Cytotoxic T protein 4	CD8 antige
55		Gene	CRP	FLT3	GATA3	IL7R	KLF1	LCK	LEF1	PLAUR	TNFSF13B	IL8	GZMB	TNFSF6	TCIRG1	PRF1	11.4	IL13	CTLA4	CD8A

Gene SEQ ID 50mer HS ACC SEQ ID RNAcDNA BY55 Instruct Miller cell receptor, instructional insuperfamily member 45.8 174.3 NM, 007053 417 OID 4460 EST He. 205159 AF 150295 417 HUB Hemoglobulin superfamily member 86 He. 155376 NM, 000518 418 HUB 23-bisphospycerate mutase 87 He. 158376 NM, 000518 418 BPOM 23-bisphospycerate mutase 87 He. 15865 NM, 000518 419 MTHPD2 Methylane tetrahydrofolate 88 He. 15865 NM, 000583 420 MTHPD2 Methylane tetrahydrofolate 88 He. 16872 NM, 000583 420 MTHPD2 Methylane tetrahydrofolate Pe. 16865 NM, 000683 420 MTHPD2 Mischondria lapha 7 90 He. 301683 AK000234 420 MDI 5573 MixAl 882 protein Pe. 102689 AK000254 426 426 OID 873 KIAA 1892 protein Pe. 102699 AK000295	55	50	45	40	35	30	25	20	15	10	5
Gene Name SEQ ID 50mer HS ACC Natural killer cell receptor, immunoglobulin superfamily member HS.81743 NM_007053 460 EST HS.205159 NM_00759 1 LS.3bisphosphoglycate mutase 87 HS.155376 NM_00058 D2 Achydrogenase (NAD+ dependent), methorylateralydrofolate 88 HS.156472 NM_00058 C40hydrogenase (NAD+ dependent) 88 HS.352018 NM_00058 NM_00058 S5 Mitochondrial solute carrier HS.300496 ANV021037 ANV021037 S6 Mitochondrial solute carrier HS.301583 ANV021037 ANV021037 A2 KANA1486 protein HS.041677 NM_000586 ANV021037 A3 KANA1486 protein HS.362018 NM_0						(continued)					
480 Instruct killer cell receptor, immunoglobulin superfamily member Hs. 81743 NM_007653 480 EST Hs. 205159 AF150295 1 2,3-bisphosphoglycerate mutase 87 Hs. 198365 NM_000518 D2 Methydrogenase (VAD+ dependent), methenylterrahydrofolate 88 Hs. 156376 NM_000536 Partydrogenase (VAD+ dependent), methenylterrahydrofolate 88 Hs. 154672 NM_000536 Partydrogenase (VAD+ dependent), methenylterrahydrofolate 88 Hs. 154672 NM_000536 Sub-family B (MDR1/TAP) Pransporter (AAD-binding cassette, sub-family B (MDR1/TAP) Hs. 352018 NM_000537 86 Mitochondrial solute carrier Hs. 300697 BC032249 Immunoglobulin heavy constant mu Hs. 300698 AK000354 73 KIAA1882 protein Hs. 30069 AK000354 73 KIAA1882 protein Hs. 301658 AK000354 73 KIAA1882 protein Hs. 301658 AK000354 74 Chemokine (C-X-C motif) receptor 4 Hs. 32401 NM_000368 74 Chemokine (C-C motif) ligand 5 <t< th=""><th>Gene</th><th>Gene Name</th><th></th><th></th><th>SEQ ID 50mer</th><th>HS</th><th></th><th>ACC</th><th>SEQ ID RN</th><th>IA/cDNA</th><th></th></t<>	Gene	Gene Name			SEQ ID 50mer	HS		ACC	SEQ ID RN	IA/cDNA	
460 EST H8.205159 AF150295 11 2.3-bisphosphoglycerate mutase 87 H8.155376 NM_000518 D2 Methylene tetrahydrofolate of ehydrogenase (NAD+ dependent), methorylterahydrofolate oyclohydrolose R8 Hs.154672 NM_006636 D2 Methylene tetrahydrofolate oyclohydrolose R8 Hs.154672 NM_00636 E3 Methylene tetrahydrolose NM_000536 NM_000536 Transporter 1, ATP-binding cassette, mutanoglobulin beavy constant mu Hs.352018 NM_000593 6 Kanyopherin alpha 6 (importin alpha 7) 90 Hs.300496 AM021037 365 Miltochondrial solute carrier Hs.300697 AW021037 73 KIAA1486 protein Hs.102669 AK000354 73 KIAA186 protein Hs.104157 AW968823 73 KIAA186 protein Hs.241392 NM_000366 73 KIAA186 protein Hs.241392 NM_000286 73 KIAA188, CC-C motif) ligand 5 Hs.241392 NM_000286 74 Chemokine (C-C-C motif) ligand 5 Hs.241392 NM_00	BY55	Natural killer ce immunoglobulir	ell receptor, n superfamily me	ember		Hs.81743		NM_007053			
11 2.3-bisphosphoglycerate mutase 86 Hs.155376 NM_000518 D2 Methylene tertahydrofolate dehydrogenase (NAD+ dependent), methenyltetrahydrofolate cyclohydrotose 88 Hs.154672 NM_0006836 B2 Methylene tertahydrofolate dehydrogenase (NAD+ dependent), methenyltetrahydrofolate cyclohydrotose 148.362018 NM_000693 B3 Transporter 1, ATP-binding cassette, sub-tamily B (MDR117AP) 90 Hs.362018 NM_000693 B6 Karyopherin alpha 6 (Importin alpha 7) 90 Hs.301553 AW021037 B5 Mitochondrial solute carrier Hs.300697 BC032249 T3 KIAA1486 protein Hs.210958 AK000354 T4 Chemokine (C-X-C motif) receptor 4 Hs.241392 NM_007865 Chemokine (C-X-C motif) ligand 5 Hs.241392 NM_00286 Chemokine (C-C motif) ligand 5 Hs.24392 NM_00286 Interleukin 5 Hs.89679 NM_00286 Interleukin 6 Hs.89679 NM_002350 Killer cellectin-like receptor subfamily 7 Hs.89687 NM_002350	OID 4460	EST			85	Hs.205159		AF150295	417		
10.2.3-bisphosphoglycerate mutase 87 Hs.198365 NM_001724 DD2 Methylene tetrahydrofolate dehydrogenase (NAD+ dependent), methylene tetrahydrofolate methydrogenase (NAD+ dependent), and then whiterhand of the control of the co	HUB	Hemoglobin, be	eta		86	Hs.155376		NM_000518	418		
D2 Methylene tetrahydrofolate dehydrogenase (NAD+ dependent), methylene tetrahydrofolate cyclohydrogenase (NAD+ dependent), methylene dehydrogenase (NAD+ dependent), methylene dehydrogenase (NAD+ dependent), methylene dehydrogenase (NAD+ dependent), methylene dehydrogenase (NAD+ dependent) Hs.352018 NM_000593 6 Karyopherin alpha 6 (importin alpha 7) 90 Hs.301553 AW021037 385 Mitochondrial solute carrier Hs.300697 BC032249 73 KIAA1486 protein Hs.210958 AB040919 73 KIAA1486 protein Hs.104157 AW968823 73 KIAA1486 protein Hs.104157 AW968823 73 KIAA1882 protein Hs.104157 AW968823 73 Chemokine (C-X-C motif) receptor 4 Hs.39414 NM 003467 74 Chemokine (C-X-C motif) ligand 5 Hs.241392 NM_002885 74 Chemokine (C-C motif) ligand 5 Hs.241392 NM_000986 75 (CANTBS, SCYA5) Hs.39879 NM_000600 76 Interfeukin 2 Hs.89679 NM_000380 8 F, member 1 Hs.183125 NM_0002850 9 <t< td=""><td>BPGM</td><td>2,3-bisphospho</td><td>oglycerate mutas</td><td>se.</td><td>87</td><td>Hs.198365</td><td></td><td>NM_001724</td><td>419</td><td></td><td></td></t<>	BPGM	2,3-bisphospho	oglycerate mutas	se.	87	Hs.198365		NM_001724	419		
formation of the composition	МТНБD2	Methylene tetra dehydrogenase methenyltetrahy cyclohydrolose	ahydrofolate (NAD+ depenc ydrofolate	lent),	88	Hs.154672		NM_006636	420		
6 Karyophein alpha 6 (importin alpha 7) 90 Hs.301553 AW021037 365 Mitochondrial solute carrier Hs.300496 AI114652 73 KilAA1486 protein Hs.300697 BC032249 73 KIAA1882 protein Hs.102669 AR00354 73 KIAA1882 protein Hs.102669 AK00354 4 Chemokine (C-X-C motif) receptor 4 Hs.104157 AW968823 5 Chemokine (C-X-C motif) receptor 4 Hs.89414 NM 003467 6 CD69 antigen (p60, early T-cell activation antigen) Hs.82401 NM_000586 7 Chemokine (C-C motif) ligand 5 Hs.241392 NM_000586 8 Interleukin 6 Hs.93913 NM 000586 9 Hiterleukin 6 Hs.93913 NM 000586 1 Killercell lectin-like receptor subfamily full member 1 Hs.183125 NM_000586 1 F, member 1 Hs.80887 NM_002350	TAP1	Transporter 1, \etasses sub-family B (N	ATP-binding cas IDR1/TAP)	sette,		Hs.352018		NM_000593			
365 Mitochondrial solute carrier Hs.300496 A114652 73 KIAA1486 protein Hs.210958 BC032249 73 KIAA1892 protein 94 Hs.10269 AK000354 73 KIAA1892 protein 94 Hs.104157 AW968823 4 Chemokine (C-X-C motif) receptor 4 Hs.89414 NM 003467 A Chemokine (C-X-C motif) ligand 5 Hs.82401 NM_001781 Chemokine (C-C motif) ligand 5 Hs.241392 NM_000600 Interleukin 6 Hr.93913 NM 000600 Interleukin 2 Hs.89679 NM 000586 F, member 1 Hs.183125 NM_000580 F, member 1 Hs.80887 NM_002350 related oncogene homolog Hs.80887 NM_002350	KPNA6	Karyopherin alp	oha 6 (importin al	pha7)	06	Hs.301553		AW021037	422		
73 KIAA1486 protein Hs.300697 BC032249 73 KIAA1486 protein 94 Hs.102669 AB040919 73 KIAA1892 protein 94 Hs.102669 AK000354 4 Chemokine (C-X-C motif) receptor 4 Hs.89414 NM 003467 CD69 artigen (p60, early T-cell activation antigen) Hs.82401 NM_001781 Chemokine (C-C motif) ligand 5 Hs.241392 NM_000585 Interleukin 6 Hs.39313 NM 000600 Interleukin 2 Hs.89679 NM_00586 F, member 1 Hs.mber 1 Hs.89679 NM_016523 F, member 1 Hs.80887 NM_002350 related oncogene homolog Hs.80887 NM_002350	OID 4365	Mitochondrial s	olute carrier			Hs.300496		AI114652			
73 KIAA1486 protein 94 Hs.102669 AB040919 73 KIAA1892 protein 94 Hs.102669 AK000354 4 Chemokine (C-X-C motif) receptor 4 Hs.89414 NM 003467 A Chemokine (C-X-C motif) receptor 4 Hs.89414 NM 003467 CD69 antigen (p60, early T-cell activation antigen) Hs.82401 NM 001781 Chemokine (C-C motif) ligand 5 Hs.241392 NM_000585 RANTBS, SCYA5) Hs.93913 NM 000600 Interleukin 6 Hs.99679 NM 000586 Interleukin 2 Hs.183125 NM_016523 F, member 1 Hs.183125 NM_000586 related oncogene homolog Hs.80887 NM_002350	IGHM	Immunoglobulir	n heavy constan	ıt mu		Hs.300697		BC032249			
73 KIAA1892 protein 94 Hs.102669 AK000354 4 EST Hs.104157 AW968823 4 Chemokine (C-X-C motif) receptor 4 activation antigen) Hs.89414 NM 003467 CD69 antigen (p60, early T-cell activation antigen) Hs.82401 NM 001781 Chemokine (C-X motif) ligand 5 (RANTBS, SCYA5) Hs.241392 NM_000585 Interleukin 6 Hs.93913 NM 000600 Interleukin 2 Hs.89679 NM 000586 Interleukin 2 Hs.183125 NM_006586 F, member 1 Hs.member 1 Hs.183125 NM_005350 v-yes-1 Yamaguchi sarcoma viral 102 Hs.80887 NM_002350	OID 573	KIAA1486 prote	uie			Hs.210958		AB040919			
4 Chemokine (C-X-C motif) receptor 4 activation antigen) Hs. 104157 AW968823 CD69 antigen (p60, early T-cell activation antigen) Hs.82401 NM_003467 CD69 antigen (p60, early T-cell activation antigen) Hs.82401 NM_001781 Chemokine (C-C motif) ligand 5 (RANTBS, SCYA5) Hs.241392 NM_0002985 Interleukin 6 Interleukin 6 Interleukin 2 Interleukin 2 Interleukin 2 Interleukin 2 Hs.89679 Hs.89679 NM_000586 I Killer cell lectin-like receptor subfamily F, member 1 F, member 1 related oncogene homolog Hs.80887 NM_002350	OID 873	KIAA1892 prote	nie		94	Hs.102669		AK000354	426		
t Chemokine (C-X-C motif) receptor 4 Hs.89414 NM 003467 CD69 antigen (p60, early T-cell activation antigen) Hs.82401 NM_001781 Chemokine (C-C motif) ligand 5 Hs.241392 NM_002985 Chemokine (C-C motif) ligand 5 Hs.93913 NM 000600 Interleukin 6 Hs.93913 NM 000600 Interleukin 2 Hs.89679 NM 000586 Killer cell lectin-like receptor subfamily F, member 1 Hs.183125 NM_00586 F, member 1 Hs.80887 NM_002350 v-yes-1 Yamaguchi sarcoma viral related oncogene homolog 102 Hs.80887 NM_002350	OID 3	EST				Hs.104157		AW968823			
CD69 antigen (p60, early T-cell activation antigen) Hs.82401 NM_001781 Chemokine (C-C motif) ligand 5 (RANTBS, SCYA5) Hs.241392 NM_002985 Interleukin 6 Interleukin 2 Hs.93913 NM 000600 Killer cell lectin-like receptor subfamily F, member 1 Hs.183125 NM_016523 v-yes-1 Yamaguchi sarcoma viral related oncogene homolog 102 Hs.80887 NM_002350	CXCR4	Chemokine (C-	X-C motif) recep	otor 4		Hs.89414		NM 003467			
Chemokine (C-C motif) ligand 5 Hs.241392 NM_002985 (RANTBS, SCYA5) Hs.93913 NM 000600 Interleukin 6 Hs.93913 NM 000600 Interleukin 2 Hs.89679 NM 000586 Killer cell lectin-like receptor subfamily F, member 1 Ho.183125 NM_016523 F, member 1 Hs.80887 NM_002350 v-yes-1 Yamaguchi sarcoma viral related oncogene homolog 102 Hs.80887 NM_002350	6900	CD69 antigen (activation antige	p60, early T-cel en)	_		Hs.82401		NM_001781			
Interleukin 6 Hs.93913 NM 000600 Interleukin 2 Hs.89679 NM 000586 Killer cell lectin-like receptor subfamily F, member 1 Hs.183125 NM_016523 V-yes-1 Yamaguchi sarcoma viral related oncogene homolog 102 Hs.80887 NM_002350	CCL5	Chemokine (C- (RANTBS, SCY	.C motif) ligand ((A5)	2		Hs.241392		NM_002985			
Killer cell lectin-like receptor subfamily101Hs.183125NM_016523F, member 1Hs.80887NM_002350v-yes-1 Yamaguchi sarcoma viral102Hs.80887NM_002350	IL6	Interleukin 6				Hs.93913		NM 000600			
Killer cell lectin-like receptor subfamily101Hs.183125NM_016523F, member 1v-yes-1 Yamaguchi sarcoma viral102Hs.80887NM_002350	1L2	Interleukin 2				Hs.89679		NM 000586			
v-yes-1 Yamaguchi sarcoma viral 102 Hs.80887 NM_002350 related oncogene homolog	KLRF1	Killercelllectin- F, member 1	likereceptorsub	family	101	Hs.183125		NM_016523	433		
	ΓΥN	v-yes-1 Yamag related oncoger	uchi sarcoma vi ne homolog	ral	102	Hs.80887		NM_002350	434		

55	50	45	40	35	30	25	20	15	10	5
					(continued)					
Gene	Gene Name			SEQ ID 50mer	HS		Acc	SEQ ID RNA/cDNA	AV/cDNA	
IL2RA	Interleukin 2 receptor, alpha	ceptor, alpha			Hs.1724		NM 000417			
CCL4	Chemokine (C-	Chemokine (C-C motif) ligand 4, SCYA4		04	Hs.75703		NM_002984	436		
OID 6207	EST				Hs.92440		D20522			
ChGn	Chondroitin beta 1,4 N-acetylgalactosami	Chondroitin beta 1,4 N-acetylgalactosaminyltransferase	Φ		Hs.11260		NM_018371			
OID 4281	EST			107	Hs.34549		AA053887	439		
CXCL9	Chemokine (C-)	Chemokine (C-X-C motif) ligand 9 (MIG)			Hs.77367		NM_002416			
CXCL10	Chemokine (C-) SCYB10	Chemokine (C-X-C motif) ligand 10, SCYB10	0,		Hs.2248		NM_001565			
IL17	Interleukin 17 (cytotoxic T-lymphocyte-associate esterase 8)	Interleukin 17 (cytotoxic Tlymphocyte-associated serine esterase 8)			Hs.41724		NM_002190			
IL15	Interleukin 15				Hs.168132		NM_000585			
IL10	Interleukin 10				Hs.193717		NM_000572 NM_			
IFNG	Interferon, gamma	ma			Hs.856		000619			
HLA-DRB1	Major histocompat class II, DR beta 1	Major histocompatibility complex, class II, DR beta 1			Hs.308026		NM_002124			
CD8B1	CD8 antigen, be	CD8 antigen, beta polypeptide 1 (p37)	37)		Hs.2299		NM_004931			
CD4	CD4 antigen (p55) 3,	55) 3,			Hs.17483		NM_000616			
CXCR3	Chemokine (C-) GPR9	Chemokine (C-X-C motif) receptor GPR9	r		Hs.198252		NM_001504			
OID 7094	XDx EST 479G12	12			NA		NA			
OID 7605	EST				Hs.109302		AA808018			
CXCL1	Chemokine (C-X-C motif) ligar (melanoma growth stimulating activity, alpha)	Chemokine (C-X-C motif) ligand 1 (melanoma growth stimulating activity, alpha)			Hs.789		NM_001511			

55	50	45	40	35	30	25	20	15	10	5
					(continued)					
Gene	Gene Name			SEQ ID 50mer	HS		ACC	SEQ ID R	SEQ ID RNA/cDNA	
OID 253	EST				Hs.83086		AK091125			
GPI	Glucose phospi	Glucose phosphate isomerase			Hs.409162		NM_000175			
CD47	CD47 antigen (integrin-associa	CD47 antigen (Rh-related antigen, integrin-associated signal transducer)	Jen, ducer)		Hs.82685		NM_001777			
HLA-F	Major histocom class I, F	Major histocompatibility complex, class I, F	,x,		Hs.377850		NM_018950			
OID 5350	EST				Hs.4283		AK055687			
TCRGC2	T cell receptor (T cell receptor gamma constant	t 2		Hs.112259		M17323			
OID 7016	EST				NA		BI018696			
PTGS2	Prostaglandin-endoperoxide synthase 2 (prostaglandin Gaynthase and cyclooxygenas	Prostaglandin-endoperoxide synthase 2 (prostaglandin G/H synthase and cyclooxygenase)			Hs.196384		NM_000963			
OID 5847	Hypothetical pri	Hypothetical protein FLJ32919			Hs.293224		NM 144588			
PRDM1	PR domain con domain	PR domain containing 1, with ZNF domain	INF		Hs.388346		NM_001198			
CKB	Creatine kinase, Brain	e, Brain			Hs.173724		NM 001823			
TNNI3	Troponin I, cardiac	Jiac			Hs.351382		NM 000363			
TNNT2	Troponin T2, cardiac	ardiac			Hs.296865		NM 000364			
MB	Myoglobin				Hs.118836		NM 005368			
SLC7A11	Solute carrier fa acid transporter 11	Solute carrier family 7, (cationic amino acid transporter, y+ system) member 11	amino ember		Hs.6682		NM_014331			
TNFRSF5	tumor necrosis factor receptor superfamily, member 5; CD40	factor receptor ember 5; CD40			Hs.25648		NM_001250			
TNFRSF7	tumor necrosis factor receptor superfamily, member 7; CD27	factor receptor ember 7; CD27			Hs.355307		NM_001242			
CD86	CD86 antigen (C B7-2 antigen)	CD86 antigen (CD28 antigen ligand 2, B7-2 antigen)	and 2,		Hs.27954		NM_175862			

55	50	45	40	35	30	20	25	20	15	10	5	
					(continued)	(pənı						
Gene	Gene Name			SEQ ID 50mer	HS			ACC	SEQ ID F	SEQ ID RNA/cDNA		
AIF1v2	Allograft inflamr variant 2	Allograft inflammatory factor 1, splice variant 2	splice		Hs.76364	4		NM_004847				
EBV BCLF-1	BCLF-1 major capsid	capsid			NA			AJ507799				
EBV	EBNA repetitive sequence	esednence			A A			AJ507799				
CMV p67	79dd				NA			X17403				
CMV TRL7	c6843-6595				A A							
CMV IE1e3	IE1 exon 3				A A			X17403				
CMV IE1e4	IE1 exon 4 (40 variants)	variants)			NA			X17403				
EBV EBNA-1	EBNA-1 cording region	g region			A A			AJ507799				
EBV BZLF-1	Zebra gene				NA			AJ507799				
EBV EBN	EBNA repetitive sequence	esdneuce			NA			AJ507799				
EBV EBNA-LP	Short EBNA leader peptide exon	ader peptide exc	uc		NA			AJ507799				
CMV IE1	IE1S				NA			X17403				
CMV IL1	IE1-MC (exon 3)	3)			NA			X17403				
CLC	Charot-Leyden crystal protein	crystal protein			Hs.889			NM 001828				
TERF2IP	telomeric repeat binding factor 2, interacting protein	ıt binding factor əin	2,		Hs.274428	28		NM_018975				
HLA-A	Major histocompatibility complex, class I, A	patibility comple	ex,		Hs.181244	44		NM_002116				
OID 5891	EST 3' end				None			AW297949				
MSCP	mitochondrial solute carrier protein	olute carrier prc	tein		Hs.283716	16		NM 018579				
DUSP5	dual specificity phosphatase	phosphatase 5			Hs.2128			NM 004419				
PRO1853	Hypothetical pro	Hypothetical protein PRO1853			Hs.433466	99		NM_018607				
OID 6420	73A7, FLJ00290 protein	0 protein			Hs.98531	1		AK090404				
CDSN	Corneodesmosin	in			Hs.507			NM 001264				
OID 4269	EST				Hs.44628	8		BM727677				

5 10 15		SEQ ID RNA/cDNA	28)46	03			9	6 4	6 4 4 305	66 44 4719	6 4 305 319	6 4 305 719 3	6 4 4 305 719 3 3	6 4 4 305 719 3 3 62	6 4 4 305 3 3 3 11 62 110	6 4 4 305 719 110 62 110	6 44 305 719 3 3 62 62 62 62 63 66 66 66 67 66 66 67 66 66 67 66 66 67 66 66	6 4 4 305 719 11 62 62 110 888 66 66 66 66 66 66 66 66 66 66 66 66	6 4 4 305 719 62 62 110 888 6 6 553	6 4 4 305 3 3 3 411 6 6 6 6 6 888 6 6 6 888 88 88 88 88 88	66 44 305 33 52 110 888 66 66 553 553
20		ACC	NM 001028	NM_002046	NM 001003	AI364926	AB023156	BE618004	NM_004805	NM_005719	BC041913	AF001542	BG260891	NM 001462	NM_022110	NM_013388	BE887646	NM_015653	NM_015435	NM_004653	X17403	NM 152312
25	(1																					
30	(continued)	SH	Hs.409158	Hs.169476	Hs.424299	ΨV	Hs.380978	Hs.381302	Hs.194638	Hs.293750	Hs.17132	Hs.356442	Hs.28310	Hs.99855	Hs.99134	Hs.279784	Hs.250824	Hs.144505	Hs.48320	Hs.80358	NA AN	Hs.86543
35		SEQ ID 50mer																				
40				phate	e, P1 1	nd 1	anger),	e 5'	NA directed)	s complex,			43932 mucin	r-like 1 FK506		ment binding	C200227	in		, ۲		J35207
45		ame	Ribosomal protein S25	Glyceraldehyde-3-phosphate dehydrogenase	Ribosomal protein, large, P1	qz23b07.x1 cDNA, 3' end /clone=IMAGE:2027701	Solute carrier family 9 (sodium/hydrogen exchanger), isoform 8	IMAGE:3865861 5 clone 5'	Polymerase (RNA) II (DNA directed) polypeptide D	Actin related protein 2/3 complex, subunit 3, 21 kDa	pu	PRO1073 protein	EST, weakly similar to A43932 mucin 2 precursor, intestinal	Formyl peptide receptor-like 1 FK506	binding protein like	prolactin regulatory element binding	Hypothetical protein LOC200227	DKFZP566F0546 protein	Ring finger protein 19	SMC (mouse) homolog, Y chromosome (SMCY)	CMV HCMVUL109	Hypothetical protein FLJ35207
50		Gene Name	Ribosom	Glyceraldehyde dehydrogenase	Ribosom	qz23b07. /clone=IN	Solute ca (sodium/h isoform 8	IMAGE:3	Polymerase (F polypeptide D	Actin rela subunit 3	EST 3' end	PRO107;	EST, wea	Formyl p	d binding p	prolactin	Hypothet	DKFZP5(Ring fing	SMC (mc	CMV HC	Hypothet
55		Gene	RPS25	GAPD	RPLP1	OID_5115	SLC9A8	OID 1512	POLR2D	ARPC3	OID 6282	PRO1073	OID_7222	FPRL1	FKBPL	PREB	OID 1551	OID 7595	RNF19	SMCY	OID 4184	OID 7504

5																						
10		SEQ ID RNA/cDNA																				
15		SEQ																				
20		ACC	NM_006260	NM_021205	NM 022752	NM_002575	NM 001428	AW297325	X17403	AL540399	AW837717	X17403	BG461987	NM 018183	086000 MN	NM_005539	BG772661	NM 001032	AW592876	X17403	NM 004665	
25																						
30	(continued)	HS	Hs.9683	Hs.20252	Hs.13323	Hs.75716	Hs.254105	Hs.438092	NA	Hs.285401	Hs.375145	NA	Hs.144814	Hs.380419	Hs.337766	Hs.124029	Hs.124675	Hs.539	Hs.352323	AN	Hs.121102	
35		SEQ ID 50mer																				
40			ubfamily C,	member U	2059	nase nin),				or, beta,	-e11 uence						otide					
45			homolog, su	gene family,	rotein FLJ2	teine) protei B (ovalbun	ha		RL2 (IRL2)	ant mRNA c ctor 2 recept anulocyte-	281299-062 ,, mRNA sec	S28			otein L18a	nosphate-5- 40kDa	ciated nucle	otein S29		L122		
50		Gene Name	DnaJ (Hsp40) homolog, subfamily C, member 3	Ras homolog gene family, member U	Hypothetical protein FLJ22059	Serine (or cysteine) proteinase inhibitor, clade B (ovalbumin), member 2	Enolase 1, alpha	EST 3' end	CMV HCMVTRL2 (IRL2)	Upstream variant mRNA of colony stimulating factor 2 receptor, beta, low-affinity (granulocytemacrophage)	CM2-LT0042-281299-062-e11 LT0042 cDNA, mRNA sequence	CMV HCMVUS28	EST	MOP-3	Ribosomal protein L18a	Inositol polyphosphate-5-phosphatase, 40kDa	Immune associated nucleotide	Ribosomal protein S29	EST 3' end	CMV HCMVUL122	vanin 2	
55		Gene	DNAJC3	ARHU	OID 7200	SERPINB2	ENO1	OID 7696	OID 4173	CSF2RB	OID_7410	OID 4180	OID 5101	MOP3	RPL18A	INPP5A	hIAN7	RPS29	OID 6008	OID 4186	VNN2	

5																			
10		SEQ ID RNA/cDNA																	
15		SEQ IC																	
20		Acc	480F8	BC038439	AW850041	NM_001402	NM 006191	NM_002046	NM_001273	NM_016521	M_021080	NM 018986	AK093608	AW452467	AI392805	NM_002964	BM467823	NM_000123	NM_001030
25											344127 Hs.								
30	(continued)	HS	NA	Hs.355841	Hs.165695	Hs.422118	Hs.374491	Hs.169476	Hs.74441		Hs.142908 Hs.344127 Hs. 61053		Hs.434526	Hs.372917	Hs.368921	Hs.416073	Hs.103804	Hs.48576	Hs.195453
35		SEQ ID 50mer																	
40						ngation	2G4, 38kDa	hate	ONAbinding	270)	sophila)	20356				otein A8	nuclear ffold	nplementing (xeroderma intation rome))	
45						inslation eld	associated 2	de-3-phosp se	iin helicase l	ein (LOC51	iolog 1 (Drc	orotein FLU				-binding pro	rogeneous itein U (scar	ir cross-con deficiency, tion group 5 , compleme kayne synd	otein S27 imulin 1)
50		Gene Name	480F8	EST	EST	Eukaryotic translation elongation factor 1 alpha 1	Proliferation-associated 2G4, 38kDa	Glyceraldehyde-3-phosphate dehydrogenase	Chromodomain helicase DNA binding protein 4	E2F-like protein (LOC51270)	Disabled homolog 1 (Drosophila)	Hypothetical protein FLJ20356	462H9 EST	EST 3' end	EST 3' end	S100 calcium-binding protein A8 (calgranulin A)	HNRPU Heterogeneous nuclear ribonucleoprotein U (scaffold attachment factor A)	Excision repair cross-complementing rodent repair deficiency, complementation group 5 (xeroderma pigmentosum, complementation group G (Cockayne syndrome))	Ribosomal protein S27 (metallopanstimulin 1)
55		Gene	OID 7057	OID 4291	OID 1366	EEF1A1	PA2G4	GAPD	CHD4	OID 7951	DAB1	OID 3406	OID 6986	OID 5962	OID 5152	S100A8	HNRPU	ERCC5	RPS27

55	50	45	40	35	30	25	20	15	10	5
					(continued)					
Gene	Gene Name			SEQ ID 50mer	HS		ACC	SEQ ID R	SEQ ID RNA/cDNA	
ACRC	acidic repeat co	acidic repeat containing (ACRC),			Hs.135167		NM 052957			
PSMD11	Proteasome (prosome, macro 26S subunit, non-ATPase, 11	Proteasome (prosome, macropain) 26S subunit, non-ATPase, 11	(u)		Hs.90744		AI684022			
OID 1016	FLJ00048 protein	nie			Hs.289034		AK024456			
OID 1309	AV706481 cDNA	Ą			None		AV706481			
OID_7582	Weakly similar	Weakly similar to ZINC FINGER					AK027866			
PROTEIN	PROTEIN 142									
OID_4317	ta73c09.x1 3' end /clone=IMAGE:20 Protein S 15	ta73c09.x1 3' end /clone=IMAGE:2049712 Ribosomal Protein S 15	nal		Hs.387179		Al318342			
OID 5889	3' end /done=Il	3' end /clone=IMAGE:3083913			Hs.255698		AW297843			
UBL1	Ubiquitin-like 1 (sentrin)	(sentrin)			Hs.81424		NM 003352			
OID 3687	EST				None		396E0W			
OID 7371	EST 5'				Hs.290874		BE730505			
SH3BGRL3	SH3 domain binding glutamic acid-rich protein like 3	nding glutamic n like 3			Hs.109051		NM_031286			
SEMA7A	Sema domain, immunogl domain (Ig), and GPI mer anchor, (semaphorin) 7A	Sema domain, immunoglobulin domain (Ig), and GPI membrane anchor, (semaphorin) 7A			Hs.24640		NM_003612			
OID 5708	EST 3' end				Hs.246494		AW081540			
OID 5992	EST 3'end				Hs.257709		AW467992			
IL21	Interleukin 21				Hs.302014		NM 021803			
HERC3	Hect domain and RLD	nd RLD 3 (HERC3)	<u>@</u>		Hs.35804		NM 014606			
OID 7799	AluJo/FLAM SINE/Alu	NE/Alu					AW837717			
P11	26 serine protease	ase			Hs.997		NM 006025			
OID 7766	EAST 3' end				Hs.437931		AW294711			

5																						
10		A/cDNA																				
15		SEQ ID RNA/cDNA																				
20		ACC	NM_012456	AJ310543	NM 003192	NM_006315	AL834168	NM_012142	X68264	NM_003173	NM 014148	AW063780	NM 004184	AL832642	AL705961	AI625119	AA136584	AK025472	NM_080475	BF968628	NM 014868	NM 013446
25																						
30	(continued)	HS.	Hs.235750	Hs.6523	Hs.75064	Hs.8834	Hs.288872	Hs.36794	AN	Hs.37936	Hs.278944	Hs.279121	Hs.82030	Hs.169610	Hs.13264	Hs.436022	Hs.70877	Hs.288156	Hs.350958	none	Hs.5094	Hs.7838
35		SEQ ID 50mer																				
40			ondrial	egans)	0 0		C	in 1		3-9			tase				clone	56766	nase iin),			1, 1
45			translocase of inner mitochondrial membrane 10 (yeast) homolog (TIMM10)	Eg1 nine homolog 1 (C. elegans)	Tubulin-specific chaperone c	rotein 3	170F9, hypothetical protein FL121439	cyclin D-type binding-protein 1 (CCNDBP1)	exons 1 &2	Suppressor of variegation 3-9 homolog 1 (Drosophila)	otein	rom T cells	Tryptophanyl-tRNA synthetase				fetal retina 937202 cDNA clone IMAGE:565899	Hypothetical protein MGC26766	Serine (or cysteine) proteinase inhibitor, clade B (ovalbumin), member 11	E:4359351 5'	rotein 10	Makorin, ring finger protein, 1
50		Gene Name	translocase o membrane 10 (TIMM10)	Eg1 nine hon	Tubulin-spec	Ring finger protein 3	170F9, hypot FL121439	cyclin D-type (CCNDBP1)	MUC18 gene exons 1 &2	Suppressor of variegatinhomolog 1 (Drosophila)	HSPC048 protein	EST 3' end from T cells	Tryptophanyl	107H8	119F12	EST 3' end	fetal retina 9372 IMAGE:565899	Hypothetical	Serine (or cys inhibitor, clad member 11	58G4, IMAGE:4359351	Ring finger protein 10	Makorin, ring
55		Gene	TIMM10	EGLN1	TBCC	RNF3	OID_6451	CCNDBP1	OID 8063	SUV39H1	HSPC048	OID 5625	WARS	OID 6823	OID 7073	OID 5339	OID_4263	MGC26766	SERPINB 11	OID 6711	RNF10	MKRN1

55	50	45	40	35	30	25	20	15	10	5
					(continued)					
Gene	Gene Name			SEQ ID 50mer	SH		ACC	SEQ ID RNA/cDNA	NA/cDNA	
RPS16	ribosomal protein S16	otein S16			Hs.397609		NM 001020			
BAZ1A	Bromodomair domain, 1A	Bromodomain adjacent to zinc finger domain, 1A	inger	Hs.8858	Hs.8858		NM_013448			
OID_5998	EST 3' end				Hs.330268		AW468459			
ATP5L	ATP synthase mitochondrial	ATP synthase, H+ transporting, mitochondrial F0 complex, subunit g	unit g		Hs.107476		NM_006476			
OID_6393	52B9				NA		52B9			
RoXaN	Ubiquitous tetratricopeptid containing protein RoXaN	Ubiquitous tetratricopeptide containing protein RoXaN			Hs.25347		BC004857			
NCBP2	Nuclear cap bi 20kDa	Nuclear cap binding protein subunit 2, 20kDa	unit2,		Hs.240770		NM_007362			
OID_6273	EST 3' end						AW294774			
HZF12	zinc finger protein 12	otein 12			Hs.158976		NM_033204			
CCL3	Chemokine ((Chemokine (C-C motif) ligand 3	~		Hs.73817		D90144			
OID_4323	IMAGE:128373 3'	73 3′			Hs.370770		AA744774			
OID_5181	tg93h12.x1 NCI_CGAP_C clone IMAGE:2116391 3' contains TAR1.t1 MER22	tg93h12.x1 NCI_CGAP_CLL1 cDNA clone IMAGE:2116391 3' similar to contains TAR1.t1 MER22	DNA ir to		NA		AI400725			
PRDX4	Peroxiredoxin 4	4 ר			Hs.83383		NM_006406			
ВТК	Bruton agamn kinase	Bruton agammaglobulinemia tyrosine kinase	osine		Hs.159494		NM_000061			
OID_6298	Importinbeta s	Importinbeta subunit mRNA			Hs.180446		A1948513			
PGK1	Phosphoglyce	Phosphoglycerate kinase 1			Hs.78771		NM_000291			
TNFRSF10A	Tumor necrosis factor rec superfamily, member 10a	Tumor necrosis factor receptor superfamily, member 10a			Hs.249190		NM_003844			
ADM	adrenomedullin	lin			Hs.394		NM_001124			
OID_357	138G5				NA		138G5			

5																			
10		SEQ ID RNA/cDNA																	
15		SEQ ID			611														
20		ACC	NM_016649	NM_015393	NM_004315	NM_012068	NM_032895	BQ022840	795290_NN	AF064257	BE502246	AI300700	AA243283	069E00 ⁻ WN	AI378046	AW063678	BF475239	C14379	NM_004064
25																			
30	(continued)	HS	Hs.88820	Hs.105460	Hs.75811	Hs.9754	Hs.417157	Hs.177376	Hs.170027	Hs.268555	Hs.122575	Hs.374836	Hs.118899	Hs.18571	Hs.309108	Hs.279116	Hs.445429	Hs.439346	Hs.238990
35		SEQ ID 50mer			279														
40			reading		drolase	or 5	376		nolog of;		tein-			ducible ndent				thetical	ibitor 1B
45			ome 20 oper	23 protein	ine amidohy se) 1	cription facto	tein MGC14		minute 2, hor itein (MDM2) nt MDM2,	slease 2	erentiation, c acid G-pro or, 4 (EDG4)	9		interferon-in d RNA deper	5	ר T cells		nilar to hypo 78	nt kinase ink
50		Gene Name	461A4 chromosome 20 open reading frame 6	DKFZP56400823 protein	N-acylsphingosine amidohydrolase (acid ceramidase) 1	Activating transcription factor 5	hypothetical protein MGC14376	EST	Mouse double minute 2, homolog of; p53-binding protein (MDM2), transcript variant MDM2,	5'-3' exoribonuclease 2	Endothelial differentiation, lysophosphatidic acid G-protein-coupled receptor, 4 (EDG4)	IMAGE:4540096	EST 3' end	Protein kinase, interferon-inducible double stranded RNA dependent activator	IMAGE:2091815	EST 3' end from T cells	EST 5'	EST Weakly similar to hypothetical protein FLJ20378	Cyclin-dependent kinase inhibitor 1B (p27, Kip1)
55		Gene	C20orf6	OID_3226	ASAH1	ATF5	OID_4887	OID_4239	MDM2	XRN2	OID_6039	OID_4210	OID_7698	PRKRA	OID_4288	OID_5620	OID_7384	OID_1209	CDKN1B

5																							
10		SEQ ID RNA/cDNA																					
15		SEQ II									634												
20		ACC	NM_000302	AK097845	AW297664	NM_012193	NM_005514	AW063921	NM_002029	NM_153437	Al392793	AW064243	469A10	AI089520	480E2	NM_013278	BM684739	AA916990	NM_007259	AA457757	AA743221	AA252909	AL832329
25																							
30	(continued)	HS	Hs.75093	Hs.283438	Hs.438118	Hs.19545	Hs.77961	Hs.279120	Hs.753	Hs.129055	Hs.160981	Hs.279139	NA	Hs.86650	NA	Hs.278911	Hs.255649	Hs.169610	Hs.6650	NA	NA	NA	Hs.294092
35		SEQ ID 50mer									302												
40			lutarate 5- xylase, ype VI)			olog 4	mplex,			n tails 2				٦,					5A (yeast)	al retina E:838756		Pu_S1	55488
45			sine, 2-oxog lysine hydro syndrome t			ophila) homo	npatibility co	om T cells	e receptor 1	ber of sperm	3' end::2107824	om T cells		hits. Aligne		O			in sorting 4	ratagene fet clone IMAG		ares_NhHM //AGE:66929	rotein LOC2
50		Gene Name	Promllagen-lysine, 2-oxoglutarate 5-dioxygenase (lysine hydroxylase, Ehlers-Danlos syndrome type VI)	EST	EST 3' end	Frizzled (Drosophila) homolog 4	Major histocompatibility complex, class I, B,	EST 3' end from T cells	Formyl peptide receptor 1	Outer dense fiber of sperm tails	tg04g01.x1 cDNA, 3' end /clone=IMAGE:2107824	EST 3' end from T cells	469A10	463C7, 4 EST hits. Aligned	480E2	Interleukin 17C	EST 3' end	CD44	Vacuolar protein sorting 45A (yeast)	aa92c03.r1 Stratagene fetal retina 937202 cDNA clone IMAGE:838756	EST	zr76a03.r1 Soares_NhHMPu_S1 cDNA clone IMAGE:669292	Hypothetical protein LOC255488
55		Gene	PLOD	OID_5128	OID_5877	FZD4	HLA-B	OID_5624	FPR1	ODF2	OID_5150	OID_5639	OID_6619	OID_6933	OID_7049	IL17C	OID_5866	CD44	VPS45A	OID_4932	OID 7821	OID_4916	OID_4891

55	50	40 45	35	30	25	20	15	10	5
				(continued)					
Gene	Gene Name		SEQ ID 50mer	HS		ACC	SEQ ID RNA/cDNA	NA/cDNA	
НАДНВ	Hydroxyacyl-Coenzyme A dehydrogenase/3-ketoacyl-Coenzyme A thiolase/enoyl-Coenzyme A hydratase (trifu protein), beta subunit,	Hydroxyacyl-Coenzyme A dehydrogenase/3-ketoacyl- Coenzyme A thiolase/enoyl- Coenzyme A hydratase (trifunctional protein), beta subunit ,		Hs.146812		NM_000183			
FLJ22757	Hypothetical pr	Hypothetical protein FLJ22757		Hs.236449		NM_024898			
RAC1	Ras-related C3 botulir substrate 1 (rho family binding protein Rac1)	Ras-related C3 botulinum toxin substrate 1 (rho family, small GTP binding protein Rac1)		Hs.173737		AK054993			
OID_6415	72D4, FLJ00290 protein	00 protein	318	Hs.98531		CA407201	650		
NM3ES1	Normalmucose 1	Normal mucosa of esophagus specific 1		Hs.112242		NM_032413			
DMBT1	Deleted in malignal transcript variant 2	Deleted in malignant brain tumors 1, transcript variant 2		Hs.279611		NM_007329			
RPS23	ribosomal protein S23	ein S23		Hs.3463		NM_001025			
ZF	HCF-binding tra Zhangfei	HCF-binding transcription factor Zhangfei		Hs.29417		NM_021212			
NFE2L3	Nuclear factor (2)-like 3	Nuclear factor (erythroid-derived 2)-like 3		Hs.22900		NM_004289			
RAD9	RAD9 homlog (S. pombe)	(S. pombe)		Hs.240457		NM_004584			
OID_6295	EST 3' end			Hs.389327		A1880607			
DEFCAP	Death effector f 4-like apoptosis variant B	Death effector filament-forming Ced- 4-like apoptosis protein, transcript variant B		Hs.104305		NM_014922			
RPL27A	Ribosomal protein L27a	ein L27a		Hs.76064		BF214146			
IL22	Interleukin 22 (IL22)	IL22)		Hs.287369		NM_020525			
PSMA4	Proteasome (pi subunit, alpha t	Proteasome (prosome, macropain) subunit, alpha type, 4, (PSMA4)		Hs.251531		NM_002789			
CCNI	cyclin I (CCNI)			Hs.79933		NM_006835			

5					
10		SEQ ID RNA/cDNA			
15		SE			
20		ACC	NM_000361	NM_006568	
25					
30	(continued)	HS	Hs.2030	Hs.59106	
35		SEQ ID 50mer HS			
40				ring finger	
45		me	modulin	Cell growth regulatory with ring finger	
50		Gene Name	Thrombomodulin	Cell grow	domain
55		Gene	ТНВD	CGR19	

5	Down Regulated			1	1										1		1			
10	Median Rank in NR																			
15	Median	4342	1775	125	7044.5	3465.5	3122.5	4153	7000.5	2732	5598.5	164.5	215.5	157	6299	2538.5	4008	4190.5	470.5	4371.5
20	Non-Para Score	779	744	735	730	730	726	725	725	722	715	710	707	703	703	702	700	700	869	269
25	SEQ ID RNA/cDNA																			
30	SEQ									722										
35	ACC	NM_001828	NM_018975	NM_002116	AW297949	NM_018579	NM_004419	109810_MN	AK090404	NM 001264	BM727677	NM_001028	NM_002046	NM_001003	AI364926	AB023156	BE618004	NM_004805	01/200_MN	BC041913
40		ystal protein	nding factor 2,	ibility complex,		te carrier	osphatase 5	in PRO1853	otein			S25	phosphate	, large, P1	, 3' end 27701	ly 9 exchanger),	clone 5'	II (DNA de D	n 2/3 complex,	
45	Gene Name	Charcot-Leyden crystal protein	telomeric repeat binding factor 2, interacting protein	Major histocompatibility compl class I, A	EST 3' end	mitochondrial solute carrier protein	dual specificity phosphatase 5	Hypothetical protein PRO1853	73A7,FLJ00290 protein	Comeodesmosin	EST	Ribosomal protein S25	Glyceraldehyde-3-phosphate dehydrogenase	Ribosomal protein, large, P1	qz23b07.x1 cDNA, 3' end /clone=IMAGE:2027701	Solute carrier family 9 (sodium/hydrogen exchanger isoform 8	IMAGE:3865861 5 clone 5'	Polymerase (RNA) II (DNA directed) polypeptide D	Actin related protein 2/3 compl subunit 3, 21kDa	EST 3' end
50																				
55	Gene	CLC	TERF2IP	HLA-A	OID_5891	MSCP	DUSP5	PRO1853	OID_6420	CDSN	OID_4269	RPS25	GAPD	RPLP1	OID_5115	SLC9A8	OID_1512	POLR2D	ARPC3	OID_6282

	Median Rank in NR Down Regulated			1.5	1.5		1	5).5	.5	1.5		5		1.5	.5		2.5	
(continued)	Non-Para Score Med	697 6754	695 6759	692 4084.5	691 1780.5	93568	689 6423	689 3882.5	5.0077 689	687 6074.5	687 6810.5	6869 989	686 3932.5	686 7584	685 2804.5	684 4690.5	684 327	683 4875.5	
	SEQ ID RNA/cDNA																		
	Acc	AF001542	B0260891	NM_001462	NM 022110	NM_013388	7 BE887646	NM_015653	NM_015435	NM_004653	X 17403	NM_152312	097900 ⁻ NN	NM_021205	NM_022752	NM_002575	NM_001428	AW297325	
	Gene Name	PRO1073 protein	EST, weakly similar to A43932 mucin 2 precursor, intestinal	Formyl peptide receptor-like 1	F506 binding protein like	Prolactin regulatory element binding	Hypothetical protein LOC200227	DKFZP566F0546 protein	Ring finger protein 19	SMC (mouse) homolog, Y chromosome (SMCY)	CMV HCMVUL109	Hypothetical protein FLJ35207	DnaJ (Hsp40) homolog, subfamily C, member 3	Ras homolog gene family, member U	Hypothetical protein FLJ22059	Serine (or cysteine) proteinase inhibitor, clade B (ovalbumin), member 2	Enolase 1, alpha	EST 3' end	
	Gene	PRO1073	OID_7222	FPRL1	FKBPL	PREB	OID_1551	OID_7595	RNF319	SMCY	OID_4184	OID_7504	DNAJC33	ARHU	OID_7200	SERPINB2	ENO1	OID_7696	(1)

194.5

671

NM_002046

Glyceraldehyde-3-phosphate

GAPD

dehydrogenase

NM_006191

Proliferation-associated 2G4,

PA2G4

38к Да

factor 1 alpha 1

4402

672

Down Regulated 5 Median Rank in NR 10 7272 4085.5 6560.5 4788.5 2620.5 6104.5 5590.5 5618.5 4838.5 107.5 4718 3753 7445 4359 6862 238 232 15 Non-Para Score 681 680 20 675 683 682 680 629 9/9 674 672 681 219 674 681 680 229 SEQ ID RNA/cDNA 25 (continued) 30 NM_005539 NM_001402 086000 MN NM 004665 **VM 014949** NM 018183 NM 001032 AW592876 AW837717 AW850041 BC038439 BG461987 BG772661 AL540399 X17403 X17403 480F8 35 ACC Upstream variant mRNA of colony Eukaryotic translation elongation stimulating factor 2 receptor, beta, LT0042 cDNA, mRNA sequence CM2-LT0042-281299-062-e 11 Immune associated nucleotide 40 Inositol polyphosphate-5low-affinity (granulocyte-Ribosomal protein L18a Ribosomal protein S29 phosphatase, 40kDa CMV HCMVUL122 CMV HCMVUS28 KIAA0907 protein macrophage) 45 **Gene Name** EST 3' end vanin 2 MOP-3 480F8 EST EST EST 50 OID_7410 OID_6008 OID_7703 OID_7057 OID 4180 OID 4186 OID_4291 OID 1366 OID 5101 CSF2RB RPL18A EEF1A1 INPP5A RPS29 MOP3 hIAN7 VNN2 Gene 55

Mon-Para Score Median Rank in NR Down Regulated Mon-Para Score Median Rank in NR Down Regulated 671		
	5199	7279.5
Non- 671 671 671 672 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668 668	667 51	667 72
SEQ ID RNA/cDNA Seq ID RNA/cDNA		
ACC NM_001273 NM_016521 NM_021080 NM_021080 NM_018986 AK093608 AW452467 Al392805 NM_0002964 NM_000123 NM_000123 NM_000123 Al684022	AK024456	AV706481
Gene Name Chromodomain helicase DNA binding protein 4 E2F-like protein (LOC51270) Disabled homolog 1 (Drosophila) Hypothetical protein FLJ20356 462H9 EST EST 3' end S100 calcium-binding protein A8 (calgranulin A) HNRPU Heterogeneous nuclear ribonucleoprotein U (scaffold attachment factor A) Excision repair cross-complementing rodent repair deficiency, complementation group 5 (xeroderma pigmentosum, complementation group G (Cockayne syndrome)) Ribosomal protein S27 (metallopaustimulin 1) acidic repeat containing (ACRC), Proteasome (prosome, macropain) 26S subunit, non-ATPase, 11	FLJ00048 protein	AV706481 cDNA
Gene CHD4 OID CHD4 OID 7951 DAB1 OID_3406 OID_5152 S100A8 HNRPU HNRPU ACRC PSMD11	OID 1016	OID 1309

Down Regulated 5 Median Rank in NR 10 1978.5 5519.5 5003.5 7751.5 5036.5 3056.5 3505.5 6224.5 5648 6499 6837 3544 310 15 Non-Para Score 20 999 999 999 999 999 665 299 664 664 664 299 665 SEQ ID RNA/cDNA 25 (continued) 30 NM_003612 NM_014606 NM_031286 AW297843 NM 003352 AW081540 AW467992 NM 021803 AW837717 BE730505 AK027866 AI318342 W03955 35 ACC Hect domain and RLD 3 (HERC3) Weakly similar to ZINC FINGER domain (lg), and GPI membrane 3' end /clone=IMAGE:3083913 Sema domain, immunoglobulin SH3 domain binding glutamic 40 anchor, (semaphorin) 7A /clone=IMAGE:2049712 Ribosomal Protein S15 Ubiquitin-like 1 (sentrin) acid-rich protein like 3 AluJo/FLAM SINE/Alu ta73c09.x13' end PROTEIN 142 Interleukin 21 45 **Gene Name** EST 3' end EST 3' end EST 5' EST 50 SH3BGRL3 OID_7582 OID_4317 OID 5889 SEMA7A 55

50		45	40	35	30 (continued)	25	20	15	10	5	
Gene Name	Gene Name			ACC	SEQ ID R	SEQ ID RNA/cDNA	Non-Para Score	Median Rank in NR	n NR	Down Regulated	
OID_6451 170F9, hyp FLJ21439	170F9, hyp FLJ21439	othe	170F9, hypothetical protein FLJ21439	AL834168			661	7126		-	
CCNDBP1 cyclin D-type (CCNDBP1)	cyclin D-typ (CCNDBP1	e bi	cyclin D-type binding-protein 1 (CCNDBP1)	NM_012142			661	1919			
OID 8063 MUC18 gene exons 1 &2	MUC18 gen	e e)	xons 1 &2	X68264			661	4692.5			
SUV39H1 Suppressor of variegation homolog 1 (Drosophila)	Suppressor homolog 1 (of \ Dro	Suppressor of variegation 3-9 homolog 1 (Drosophila)	NM_003173			661	5103		_	
HSPC048 protein	HSPC048 p	rote	nie	NM 014148			099	5981.5			
OID 5625 EST 3' end fromT cells	EST 3' end	fron	nT cells	AW063780			099	4437		_	
WARS Tryptophan	Tryptophan	yl-tF	Tryptophanyl-tRNA synthetase	NM_004184			099	905.5			
OID 6823 107H8	107H8			AL832642			629	2619			
OID 7073 119F12	119F12			AL705961			629	6837.5			
OID 5339 EST 3' end	EST 3' en	þ		AI625119			658	4414.5		1	
OID_4263 fetal retina 9372 IMAGE:565899	fetal retin IMAGE:5	a 937; 65899	fetal retina 937202 cDNA clone IMAGE:565899	AA136584			658	5870			
MGC26766 Hypotheti	Hypothetio	cal pro	Hypothetical protein MGC26766	AK025472			658	1892.5			
SERPINB1 1 Serine (or c inhibitor, cla	Serine (or inhibitor, c member 1	cyste lade l	Serine (or cysteine) proteinase inhibitor, clade B (ovalbumin), member 11	NM_080475			658	7535.5		1	
OID 6711 58G4, IMAGE:4359351	58G4, IM/	\GE:4	4359351 5'	BF968628			658	7264			
RNF10 Ring finder protein 10	Ring finde	r prot	tein 10	NM .014868			658	3127.5			
MKRN1 Makorin, r	Makorin, r	ing fir	Makorin, ring finger protein, 1	NM 013446			658	2228.5			
RPS16 ribosomal protein S16	ribosomal	prote	ein S16	NM 001020			657	165.5			
BAZ1A Bromodomain adji finger domain, 1A	Bromodoı finger dor	main a nain, 1	Bromodomain adjacent to zinc finger domain, 1A	NM_013448			657	2533			
OID 5998 EST 3' end	EST 3' en	þ		AW468459			657	6339.5			

6187.5

653

NM 015393

DKFZP56400823 protein

OID 3226

reading frame 6

Down Regulated 5 Median Rank in NR 10 5498.5 4715.5 6406.5 3397.5 2433.5 2059.5 4897.5 5427.5 7420.5 4666.5 4910 1155 7378 4838 2358 4235 6343 15 Non-Para Score 20 929 929 655 655 655 655 655 655 657 929 656 654 657 929 654 654 654 SEQ ID RNA/cDNA 25 (continued) 30 NM 006476 NM_003844 NM_016649 NM_007362 NM 006406 NM 033204 NM_000061 AW294774 NM 000291 **VM 001124** AA744774 BC004857 AI948513 AI400725 138G5 35 52B9 ACC Chemokine (C-C motif) ligand 3 ATP synthase, H+ transporting, tg93h12.x1 NCI_CGAP_CLL1 cDNA clone IMAGE:2116391 3' Tumor necrosis factor receptor Bruton agammaglobulinemia 461A4 chromosome 20 open Importin beta subunit mRNA 40 Ubiquitous tetratricopeptide Nuclear cap binding protein similar to contains TAR1.t1 Phosphoglycerate kinase I mitochondrial F0 complex, containing protein RoXaN superfamily, member 10a zinc finger protein 12 IMAGE:1283731 3' subunit 2, 20kDa Peroxiredoxin 4 adrenomedullin tyrosine kinase 45 **Gene Name** EST 3' end subunit g MER22 138G5 52B9 50 TNFRSF10 A OID 6273 OID_5181 OID 6298 OID 6393 OID 4323 OID 357 C20orf6 NCBP2 PRDX4 RoXaN HZF12 ATP5L CCL3 Gene PGK1 ADM 55

	Down Regulated			1						←					_	
	Median Rank in NR	1003	4545.5	2310	.2774.5	.4342	6.9689	5147	1330.5	7432.5	3512.5	6401.5	6400	6875	1356.5	4272.5
	Non-Para Score	653	653	653	652	652	652	652	652	652	652	651	651	651	651	650
(continued)	SEQ ID RNA/cDNA	611														
)))	ACC	NM_004315	NM_012068W4	NM_032895	BQ022840	NM_002392	AF064257	BE502246	AI300700	AA243283	NM_003690	AI378046	AW063678	BF475239	C14379	
	Gene Name	N-acylsphingosine amidohydrolase (acid ceramidase) 1	Activating transcription factor 5	hypothetical protein MGC14376	EST	Mouse double minute 2, homolog of; p53-binding protein (MDM2), transcript variant MDM2,	5'-3' exoribonuclease 2	Endothelial differentiation, lysophosphatidic acid G-protein- coupled receptor, 4 (EDG4)	IMAGE:4540096	EST 3' end	Protein kinase, interferon-inducible double stranded RNA dependent activator	IMAGE:2491815	EST 3' end from T cells	BST 5'	EST Weakly similar to hypothetical protein FLJ20378	Cyclin-dependent kinase inhibitor 1B (p27, Kipl1)
	Gene	ASAH1	ATF5	OID_4887	OID 4239	.МDM2	XRN2	OID_6039	OID 4210	OID 7698	PRKRA	OID 4288	OID 5620	OID 7384	OID_1209	CDKN1B
		279														

5		Down Regulated			1							1	1	1			1			1	
10 15		Median Rank in NR	3101	6476	6864.5	5816	229	7812.5	1156.5	4982.5	7638	6805	7110	6880.5	7128.5	6411.5	6532	4758	3371	6057	7507
20		Non-Para Score	920	9 059	9 059	9 2	650	649	649	649	649	648 6	647	647	647	647	647	646	646	646	645 7
25 30	(continued)	SEQ ID RNA/cDNA																			
35	(00)	ACC	NM_000302	AK097845	AW297664	NM_012193	NM_005514	AW063921	NM 002029	NM_153437	AI392793	AW064243	469A10	AI089520	480E2	NM 013278	BM684739	AA916990	NM_007259	AA457757	AA743221
40			Procollagen-lysine, 2- oxoglutarate 5-dioxygenase (lysine hydroxylase, Ehlers- Danlos syndrome type VI)			Frizzled (Drosophila) homolog 4	Major histocompatibility complex, class I, B	om T cells	e receptor l	Outer dense fiber of sperm tails 2	JNA, 3' end E:2107824	om T cells		hits. Aligned		0			Vacuolar protein sorting 45A (yeast)	aa92c03.r1 Stratagene fetal retina 937202 cDNA clone IMAGE:838756	
45		Gene Name	Procollagen-lysine, 2- oxoglutarate 5-dioxygenase (lysine hydroxylase, Ehlers- Danlos syndrome type VI)	EST	EST 3' end	Frizzled (Dros	Major histocor class I, B	EST 3' end from T cells	Formyl peptide receptor l	Outer dense fi	tg04g01.x1 cDNA, 3' end /clone=IMAGE:2107824	EST 3' end from T cells	469A10	463C7, 4 EST hits. Aligned	480E2	Interleukin 17C	EST 3' end	CD44	Vacuolar prote (yeast)	aa92c03.r1 Stratagene fets retina 937202 cDNA clone IMAGE:838756	EST
55		Gene	PLOD	OID 5128	OID 5877	FZD4	HLA-B	OID 5624	FPR1	ODF2	OID_5150	OID 5639	OID 6619	OID 6933	OID 7049	IL17C	OID 5866	CD44	VPS45A	OID_4932	OID 7821
									FPR1		302										

5		Down Regulated	1			_			_								1
10 15		Median Rank in NR	6962.5	6148.5	3212.5	1965.5	1533	4881	.6217	7284	219.5	4069	3378	6453	7493.5	3059	6571
20		Non-Para Score	645	645	645	644	644	644	644		643	643	643	643	643	643	642
25 30	(continued)	SEQ ID RNA/cDNA			645			650									
35	55)	ACC	AA252909	AL832329	NM_000183	NM 024898	AK054993	CA407201		NM_007329	NM 001025	NM_021212	NM_004289	NM 004584	AI880607	NM_014922	BF214146
40			ares_NhHMPu_S1 IAGE:669292	Hypothetical protein LOC255488	oenzyme A e/3-ketoacyl- hiolase/enoyl- ydratase rotein), beta	Hypothetical protein FLJ22757	Ras-related C3 botulinum toxin substrate 1 (rho family, small GTP binding protein Rac1)	90 protein	Normal mucosa of esophagus specific 1	Deleted in malignant brain tumors 1, transcript variant 2	ein S23	HCF-binding transcription factor Zhangfei	Nuclear factor (erythroid-derived 2)-like 3	g (S. pombe)		Death effector filament-forming Ced-4-like apoptosis protein, transcript variant B	tein L27a
45 50		Gene Name	zr76a03.r1 Soares_NhHMPu cDNA clone IMAGE:669292	Hypothetical pr	Hydroxyacyl-Coenzyme A dehydrogenase/3-ketoacyl-Coenzyme A thiolase/enoyl-Coenzyme A hydratase (trifunctional protein), beta subunit	Hypothetical pr	Ras-related C3 botulii substrate 1 (rho family binding protein Rac1)	72D4, FLJ00290 protein	Normal mucos specific 1	Deleted in malignant b 1, transcript variant 2	ribosomal protein S23	HCF-binding tr Zhangfei	Nuclear factor 2)-like 3	RAD9 homolog (S. pombe)	EST 3' end	Death effector filament-formii Ced-4-like apoptosis protein, transcript variant B	Ribosomal protein L27a
55		Gene	OID_4916	OID_4891	НАДНВ	FLJ22757 '	RAC1	OID 6415	NMES1	DMBT1	RPS23	ZF	NFE2L3	RAD9	OID 6295	DEFCAP	RPL27A
								318									

5		Down Regulated	_						
10 15		Median Rank in NR Down Regulated	3891	1934.5		980.5	4732.5	5510	
20		Non-Para Score N	642 3	641		641	640	640 5	
25 30	(continued)	SEQ ID RNA/cDNA Non-Para Score							
35	uoo)	ACC	NM 020525	NM_002789		NM 006835	NM 000361	NM_006568	
40			(1L22)	rosome,	macropain) subunit, alpha type, 4, (PSMA4)		lin	Cell growth regulatory with ring	
45 50		Gene Name	Interleukin 22 (IL22)	Proteasome (prosome,	macropain) sub (PSMA4)	cyclin I (CCNI)	Thrombomodulin	Cell growth reg	
55		Gene	1L22	PSMA4		CCNI	ТНВD	CGR19	
		l	l	l		1	l	l	

Gene	SEQ ID 50mer	SEQ ID RNA/cDNA	PCR forward Primer 1 SEQ ID	PCR Reverse Primer 1 SEQ ID	PCR Probe1 SEQ ID	PCR forward Primer 2 SEQ ID	PCR Reverse Primer 2 SEQ ID	PCR Probe 2 SEQ ID
HBB	36	368	700	1031	1362	1677	1925	2173
ITGB7	49	381	713	1044	1375			
TNFSF6	77	409	741	1072	1403			
TCIRG1	78	410	742	1073	1404			
PRF1	79	411	743	1074	1405			
IL4	80	412	744	1075	1406			
OID 4460	85	417	749	1080	1411			
HBB	86	418	750	1081	1412			
BPGM	87	419	751	1082	1413			
MTHFD2	88	420	752	1083	1414			
OID 4365	91	423	755	1086	1417			
OID 873	94	426	758	1089	1420			
KLRF1	101	433	765	1096	1427			
LYN	102	434	766	1097	1428			
CCL4	104	436	768	1099	1430			
OID 4281	107	439	771	1102	1433			
ASAH1	279	611	942	1273	1604	1850	2098	2346
OID 5150	302	634	965	1296	1627	1873	2121	2369
OID 6415	318	650	981	1312	1643	1889	2137	2385

Gene	Gene Name	SEQ ID 50mer	SEQ ID RNA/cDNA	n	Non-parametric Odds ratio	Fisher p-value	t-test p-value
HBB	Hemoglobin, beta		418	55	8.33	0.00	0.00
OID_4365	Mitochondrial solute carrier			53	6.16	0.00	0.00
OID 873	KIAA1892 protein		426	55	5.09	0.01	0.01
IL4	Interleukin 4			46	4.90	0.02	0.01
OID 4281	EST		439	56	5.19	0.01	0.01
IGHM	Immunoglobulin heavy constant mu			52	2.89	0.09	0.01
BPGM	2,3-bisphosphoglycerate mutase		419	43	7.31	0.01	0.01
CTLA4	Cytotoxic T-lymphocyte associated protein 4			52	1.84		0.02
SLC7A11	Solute carrier family 7, (cationic amino acid transporter, y+ system) member 11			48	2.50	0.15	0.03

5	Gene	Gene Name	SEQ ID 50mer	SEQ ID RNA/cDNA	n	Non-parametric Odds ratio	Fisher p-value	t-test p-value
	IL13	Interleukin 13			29	4.95	0.07	0.04
	OID 6207	EST			37	3.58	0.10	0.04
10	PRDM1	PR domain containing with ZNF domain			57	1.44		0.07
	LYN	v-yes-1 Yamaguchi sarcoma viral related oncogene homolog		434	55	1.08		0.08
15	KPNA6	Karyopherin alpha 6 (importin alpha 7)		422	51	1.50		0.09
	OID 7094	XDxEST479G12			35	1.13		0.09
	IL15	Interleukin 15			51	3.78	0.05	0.09
20	OID 4460	EST		417	47	2.73	0.14	0.10
	OID 7016	EST			53	2.14	0.27	0.10
25	MTHFD2	Methylene tetrahydrofolate dehydrogenase (NAD+ dependent), methenyltetrahydrofolate cyclohydrolase			43	3.50	0.07	0.11
30	TCIRG1	T-cell, immune regulator 1, ATPase, H+ transporting, lysosomal V0 protein a isoform 3	78	410	57	1.08		0.11
0.5	OID_5847	Hypothetical protein FLJ32919			45	1.08		0.12
35	CXCR4	Chemokine (C-X-C motif			56	1.29		0.12
	CXCR3	Chemokine (C-X-C motif			54	2.10	0.27	0.12
	GPI	Glucose phosphate			57	1.44	0.60	0.12
40	KLRF1	Killer cell lectin-like rece	101	433	50	1.68		0.13
	CCL5	Chemokine (C-C motif) 1			34	1.96		0.13
	CD47	CD47 antigen (Rh-related			55	1.45		0.13
45	IL10	Interleukin 10			33	1.43		0.13
70	OID 253	EST			26	1.93		0.15
	CXCL10	Chemokine (C-X-C	motif		53	1.75		0.16
	IFNG	Interferon, gamma			41	1.33		0.16
50	PRF1	Perforin 1 (pore forming	79	411	48	1.20		0.17
	IL2	Interleukin 2			33	2.00		0.17
	HLA-DRB1	Major histocompatibility			42	1.50		0.18
55	IL6	Interleukin 6			49	1.33		0.18
	IL2RA	Interleukin 2 receptor, alpha			39	2.03	0.34	0.19
	OID 573	KIAA1486 protein			8	3.00		0.19

5	Gene	Gene Name	SEQ ID 50mer	SEQ ID RNA/cDNA	n	Non-parametric Odds ratio	Fisher p-value	t-test
	CXCL9	Chemokine (C-X-C motif) ligand 9 (MIG)			46	1.71	•	0.20
	OID 3	EST			49	2.19		0.20
10	CD8B1	CD8 antigen, beta polypeptide 1 (p37)			55	1.21		0.22
	CD69	CD69 antigen (p60, early T-cell activation antigen)			30	1.71		0.23
15	OID 7605	EST			47	3.11	0.08	0.24
	TNFSF6	Tumor necrosis factor (ligand) superfamily, member 6	77	409	54	1.36		0.25
20	CXCL1	Chemokine (C-X-C motif) ligand 1 (melanoma growth stimulating activity, alpha)			20	2.00		0.26
	OID 5350	EST			49	2.08	0.26	0.28
25	CD8A	CD8 antigen, alpha polypeptide (p32)			57	1.39		0.28
	CD4	CD4 antigen (p55)			55	1.64		0.28
30	PTGS2	Prostaglandin-endoperoxide synthase 2 (prostaglandin G/H synthase and cyclooxygenase)			46	2.05	0.37	0.29
35	GZMB	Granzyme B (granzyme 2, cytotoxic T-lymphocyte-associated serine esterase 1)			40	1.81		0.33
	CCL4	Chemokine (C-Cmotif) ligand 4, SCYA4			53	2.25		0.35
40	ChGn	Chondroitin beta 1,4 N-acetylgalactosaminyltran sferase			31	2.57		0.36
45	TCRGC2	T cell receptor gamma constant 2			52	1.33		0.39
	HLA-F	Major histocompatibility complex, class I, F			54	2.36	0.17	0.40
50	TAP1	Transporter 1, ATP-binding cassette, sub-family B (MDR1/TAP)			36	1.93		0.45
	BY55	Natural killer cell receptor, immunoglobulin superfamily member			52	2.49	0.16	0.48
55	IL8	Interleukin 8			49	2.10	0.26	0.49

	Gene	ACC	SEQ ID 50mer	SEQ ID RNA/cDNA	Ref Seq Peptide Accession #	SEQ ID Protein
5	ACTB	NM_001101			NP 001092	
	GUSB	NM_000181			NP_000172	
	B2M	NM_004048			NP_004039	
10	TSN	NM_004622 INM			NP_004613	
	CCR7	NM_001838			NP 001829	
	IL1R2	NM_004633			NP_004624	
	AIF-1	NM_004847			NP_004838	
15	ALAS2	NM_000032.1			NP 000023	
	APELIN	NM_017413			NP 059109	
	CD80	NM_005191			NP_005182	
20	EPB41	NM_004437			NP_004428	
	CBLB	NM_004351			NP_733762	
	CCR5	NM_ 000579			NP 000570	
25	MME	NM_000902			NP 000893	
25	KLRC1	NM_002259			NP 002250	
	FCGR3A	NM_000569			NP 000560	
	FCGR3B	NM_000570			NP 000561	
30	LAG3	NM_002286			NP 002277	
	PECAM1	NM_000442			NP 000433	
	CD34	NM_001773			NP 001764	
35	FCGR1A	NM_000566			NP_000557	
	TFRC	NM_003234			NP_003225	
	CMA1	NM_001836			NP 001827	
	KIT	NM_000222			NP 000213	
40	MPL	NM_005373			NP 005364	
	EphB6	NM_004445			NP 004436	
	EPO-R	NM_000121.2			NP 000112	
45	Foxp3	NM_014009			NP 054728	
	GATA-1	NM_002049			NP 002040	
	ITGA2B	NM_000419			NP 000410	
	GNLY	NM_006433			NP 006424	
50	GZMA	NM_006144			NP 006135	
	НВА	NM_000558.3			NP 000549	
	HBZ	NM_005332.2			NP_005323	
55	HBD	NM_000519.2			NP_000510	
	HBE	NM_005330			NP_005321	
	HBG	NM_000559.2			NP_000550	

5	Gene	ACC	SEQ ID 50mer	SEQ ID RNA/cDNA	Ref Seq Peptide Accession #	SEQ ID Protein
	HBQ	NM_005331			NP 005322	
	HLA-DP	NM_033554			NP 291032	
	HLA-DQ	NM_002122			NP 002113	
10	ICOS	NM_012092			NP 036224	
	IL18	NM_001562			NP_001553	
	IL3	NM_000588			NP_000579	
15	ITGA4	NM_000885			NP 000876	
	ITGAM	NM_000632			NP_000623	
	ITGB7	NM_000889	49	381	NP 000880	
20	СЕВРВ	NM_005194			NP_005185	
20	NF-E2	NM_006163			NP_006154	
	PDCD1	NM_005018			NP 005009	
	PF4	NM_002619			NP 002610	
25	PRKCQ	NM_006257.1			NP 006248	
	PPARGC1	NM_013261			NP 037393	
	RAG1	NM_000448			NP 000439	
30	RAG2	NM_000536			NP 000527	
30	CXCL12	NM_000609			NP_000600	
	TNFRSF4	NM_003327			NP_003318	
	TNFSF4	NM_003326			NP_003317	
35	TPS1	NM_003293			NP_003284	
	ADA	NM_000022			NP_000013	
	СРМ	NM_001874.1			NP_001865	
40	CSF2	NM_000758.2			NP_000749	
	CSF3	NM_172219			NP_757373	
	CRP	NM_000567.1			NP_000558	
	FLT3	NM_004119			NP_004110	
45	GATA3	NM_002051.1			NP_002042	
	IL7R	NM_002185.1			NP_002176	
	KLF1	NM_006563.1			NP_006554	
50	LCK	NM_005356.2			NP_005347	
	LEF1	NM_016269.2			NP_057353	
	PLAUR	NM_002659.1			NP_002650	
	TNFSF13B	NM_006573.3			NP_006564	
55	IL8	NM_000584			NP_000575	
	GZMB	NM_004131			NP_004122	

5	Gene	ACC	SEQ ID 50mer	SEQ ID RNA/cDNA	Ref Seq Peptide Accession #	SEQ ID Protein
	TNFSF6	NM_000639	77		NP_000630	2473
	TCIRG1	NM_006019	78		NP_006010	2474
40	PRF1	NM_005041	79		NP_005032	2475
10	IL4	NM_000589	80		NP_000580	2476
	IL13	NM_002188			NP_002179	
	CTLA4	NM_005214			NP_005205	
15	CD8A	NM_001768			NP_001759	
	BY55	NM_007053			NP_008984	
	НВВ	NM_000518			NP_000509	2481
20	BPGM	NM_001724	87		NP_001715	2482
20	MTHFD2	NM_006636	88		NP_006627	2483
	TAP1	NM_000593			NP_000584	
	OID 873	AK_000354	94		NP_056212	2485
25	CXCR4	NM_003467			NP_003458	
	CD69	NM_001781			NP_001772	
	CCL5	NM_002985			NP_002976	
30	IL6	NM_000600			NP_000591	
	IL2	NM_000586			NP 000577	
	KLRF1	NM_016523	101		NP_057607	2491
	LYN	NM_002350	102		NP_002341	2492
35	IL2RA	NM_000417			NP_000408	
	CCL4	NM_002984			NP_002975	
	ChGn	NM_018371			NP_060841	
40	CXCL9	NM_002416			NP_002407	
	CXCL10	NM_001565			NP_001556	
	IL17	NM_002190			NP_002181	
	IL15	NM_000585			NP_000576	
45	IL10	NM_000572			NP_000563	
	IFNG	NM_000619			NP_000610	
	HLA-DRB1	NM_002124			NP_002115	
50	CD8B1	NM_004931			NP_004922	
	CD4	NM_000616			NP_000607	
	CXCR3	NM_001504			NP_001495	
	CXCL1	NM_001511			NP_001502	
55	GPI	NM_000175			NP_000166	
	CD47	NM_001777			NP_001768	

5	Gene	ACC	SEQ ID 50mer	SEQ ID RNA/cDNA	Ref Seq Peptide Accession #	SEQ ID Protein
	HLA-F	NM_018950			NP_061823	
	PTGS2	NM_000963			NP_000954	
	OID 5847	NM_144588			NP_653189	
10	PRDM 1	NM_001198			NP_001189	
	СКВ	NM_001823			NP_001814	
	TNNI3	NM_000363			NP_000354	
15	TNNT2	NM_000364			NP_000355	
	MB	NM_005368			NP_005359	
	SLC7A11	NM_014331			NP_055146	
00	TNFRSF5	NM_001250			NP_001241	
20	TNFRSF7	NM_001242			NP_001233	
	CD86	NM_175862			NP_787058	
	AIF1v2	NM_004847			NP_004838	
25	CMV IE1e3	NC_001347,compl			NP_040060	
	CMV IE1e4	NC_001347, compl			NP_040060	
	EV EBNA-1	NC_001345,10795			NP_039875	
30	EV BZLF-1	NC_001345, compl			NP_039871	
30	CMV IB1	NC_001347, compl			NP_040060	
	CMV IE1	NC_001347, compl			NP_040060	
	CLC	NM_001828			NP_001819	
35	TERF2IP	NM_018975			NP_061848	
	HLA-A	NM_002116			NP_002107	
	MSCP	NM_018579			NP_061049	
40	DUSP5	NM_004419			NP_004410	
	PRO1853	NM_018607			NP_061077	
	CDSN	NM_001264			NP_001255	
	RPS25	NM_001028			NP_001019	
45	GAPD	NM_002046			NP_002037	
	RPLP1	NM_001003			NP_000994	
50	POLR2D	NM_004805			NP_004796	
	ARPC3	NM_005719			NP_005710	
	FPRL1	NM_001462			NP_001453	
	FKBPL	NM_022110			NP_071393	
	PREB	NM_013388			NP_037520	
55	OID_7595	NM_015653			NP_056468	
	RNF19	NM_015435			NP_056250	

5	Gene	ACC	SEQ ID 50mer	SEQ ID RNA/cDNA	Ref Seq Peptide Accession #	SEQ ID Protein
	SMCY	NM_004653			NP_004644	
	OID_7504	NM_152312			NP_689525	
40	DNAJC3	NM_006260			NP_006251	
10	ARHU	NM_021205			NP_067028	
	OID_7200	NM_022752			NP_073589	
	SERPINB2	NM_002575			NP_002566	
15	ENO1	NM_001428			NP_001419	
	MOP3	NM_018183			NP_060653	
	RPL18A	NM_000980			NP_000971	
20	INPP5A	NM_005539			NP_005530	
20	RPS29	NM_001032			NP_001023	
	VNN2	NM_004665			NP_004656	
	OID_7703	NM_014949			NP_055764	
25	EEF1A1	NM_001402			NP_001393	
	PA2G4	NM_006191			NP_006182	
	GAPD	NM_002046			NP_002037	
30	CHD4	NM_001273			NP_001264	
	OID_7951	NM_016521			NP_057605	
	DAB1	NM_021080			NP_066566	
	OID_3406	NM_018986			NP_061859	
35	S100A8	NM_002964			NP_002955	
	ERCC5	NM_000123			NP_000114	
	RPS27	NM_001030			NP_001021	
40	ACRC	NM_052957			NP_443189	
	UBL1	NM_003352			NP_003343	
	SH3BGRL3	NM_031286			NP_112576	
	SEMA7A	NM_003612			NP_003603	
45	IL21	NM_021803			NP_068575	
	HERC3	NM_014606			NP_055421	
50	P11	NM_006025			NP_006016	
	TIMM10	NM_012456			NP_036588	
	EGLN1	AJ310543			NP_071334	
	TBCC	NM_003192			NP_003183	
	RNF3	NM_006315			NP_006306	
55	CCNDBP1	NM_012142			NP_036274	
	SUV39H1	NM_003173			NP_003164	

5	Gene	ACC	SEQ ID 50mer	SEQ ID RNA/cDNA	Ref Seq Peptide Accession #	SEQ ID Protein
	HSPC048	NM_014148			NP_054867	
	WARS	NM_004184			NP_004175	
40	SERPINB11	NM_080475			NP_536723	
10	RNF 10	NM_014868			NP_055683	
	MKRN1	NM_013446			NP_038474	
	RPS16	NM_001020			NP_001011	
15	BAZ1A	NM_013448			NP_038476	
	ATP5L	NM_006476			NP_006467	
	NCBP2	NM_007362			NP_031388	
20	HZF12	NM_033204			NP_149981	
20	CCL3	D90144			NP_002974	
	PRDX4	NM_006406			NP_006397	
	ВТК	NM_000061			NP_000052	
25	PGK1	NM_000291			NP_000282	
	TNFRSF10A	NM_003844			NP_003835	
	ADM	NM_001124			NP_001115	
30	C20orf6	NM_016649			NP_057733	
	OID_3226	NM_015393			NP_056208	
	ASAH1	NM_004315	279	611	NP_004306	
	ATF5	NM_012068			NP_036200	
35	OID_4887	NM_032895			NP_116284	
	MDM2	NM_002392			NP_002383	
	XRN2	AF064257			NP_036387	
40	PRKRA	NM_003690			NP_003681	
	CDKN1B	NM_004064			NP_004055	
	PLOD	NM_000302			NP_000293	
	FZD4	NM_012193			NP_036325	
45	HLA-B	NM_005514			NP_005505	
	FPR1	NM_002029			NP_002020	
50	ODF2	NM_153437			NP_702915	
	IL17C	NM_013278			NP_037410	
	VPS45A	NM_007259			NP_009190	
	HADHB	NM_000183			NP_000174	
	FLJ22757	NM_024898			NP_079174	
55	NMES 1	NM_032413			NP_115789	
	DMBT1	NM_007329			NP_015568	

5	Gene	ACC	SEQ ID 50mer	SEQ ID RNA/cDNA	Ref Seq Peptide Accession #	SEQ ID Protein
	RPS23	NM_001025			NP_001016	
	ZF	NM_021212			NP_067035	
	NFE2L3	NM_004289			NP_004280	
10	RAD9	NM_004584			NP_004575	
	DEFCAP	NM_014922			NP_055737	
	IL22	NM_020525			NP_065386	
15	PSMA4	NM_002789			NP_002780	
	CCNI	NM_006835			NP_006826	
	THBD	NM_000361			NP_000352	
20	CGR19	NM_006568			NP_006559	
20	HSRRN18S	X03205				
	НВВ	NG_000007				
	HLA-DRB					
25	OID_4460	AF150295		417		
	KPNA6	AW021037		422		
	OID_4365	Al114652				
30	IGHM	BC032249				
30	OID_573	AB040919				
	OID_3	AW968823				
	OID_6207	D20522				
35	OID_4281	AA053887		439		
	OID_7094					
	OID_7605	AA808018				
40	OID_253	AK091125				
	OID_5350	AK055687				
	TCRGC2	M17323				
	OID_7016	BI018696				
45	EV EBV					
	CMV p67	NC_001347				
50	CMV TRL7					
	EV EBN					
	EV EBNA-LP					
	OID_5891	AW297949				
	OID_6420	AK090404				
55	OID_4269	BM727677				
	OID_5115	Al364926				

5	Gene	ACC	SEQ ID 50mer	SEQ ID RNA/cDNA	Ref Seq Peptide Accession #	SEQ ID Protein
5	SLC9A8	AB023156	Joiner	KNA/CDNA	Accession #	Fioteni
	OID_1512	BE618004				
		BC041913				
10	OID_6282					
	PRO1073	AF001542				
	OID_7222	BG260891				
	OID_1551	BE887646				
15	OID_4184	X17403				
	OID_7696	AW297325				
	OID_4173	X17403				
20	CSF2RB	AL540399				
	OID_7410	AW837717				
	OID_4180	X17403				
	OID_5101	BG461987				
25	hIAN7	BG772661				
	OID_6008	AW592876				
	OID_4186	X17403				
30	OID_7057	480F8				
00	OID_4291	BC038439				
	OID_1366	AW850041				
	OID_6986	AK093608				
35	OID_5962	AW452467				
	OID_5152	Al392805				
	HNRPU	BM467823				
40	PSMD11	AI684022				
	OID_1016	AK024456				
	OID_1309	AV706481				
	OID_7582	AK027866				
45	OID_4317	A1318342				
	OID_5889	AW297843				
50	OID_3687	W03955				
	OID_7371	BE730505				
	OID_5708	AW081540				
	OID_5992	AW467992				
	OID_7799	AW837717				
55	OID_7766	AW294711				
	OID_6451	AL834168				

5	Gene	ACC	SEQ ID 50mer	SEQ ID RNA/cDNA	Ref Seq Peptide Accession #	SEQ ID Protein
Ö	OID_8063	X68264		, , , , , , , , , , , , , , , , , ,	71000001011 11	
	OID_5625	AW63780				
	OID_6823	AL832642				
10	OID_7073	AL705961				
	OID_5339	AI625119				
	OID_3339	AA136584				
15	MGC26766	AK025472				
10	OID_6711	BF968628				
	OID_5998	AW468459				
	OID_5998 OID_6393	52B9				
20	RoXaN	BC004857				
	OID_6273	AW294774				
	OID_6273	AV294774 AA744774				
25	OID_4323	A1400725				
20		A1948513				
	OID_6298					
	OID_357	138G5				
30	OID_4239	BQ022840				
	OID_6039	BE502246				
	OID_4210	A1300700				
25	OID_7698	AA243283				
35	OID_4288	Al378046				
	OID_5620	AW063678				
	OID_7384	BF475239				
40	OID_1209	C14379				
	OID_5128	AK097845				
	OID_5877	AW297664				
45	OID_5624	AW063921				
45	OID_5150	Al392793	302	634		
	OID_5639	AW064243				
50	OID_6619	469A10				
	OID_6933	AI089520				
	OID_7049	480E2				
	OID_5866	BM684739				
	CD44	AA916990				
55	4932	AA457757				
	OID_7821	AA743221				

(continued)

Ref Seq **SEQ ID SEQ ID** Peptide SEQ ID ACC RNA/cDNA Accession # Protein Gene 50mer OID_4916 AA252909 OID_4891 AL832329 RACI AK054993 OID_6415 CA407201 318 650 OID_6295 AI880607 RPL27A BF214146

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Table 3: Viral genomes were used to design oligonucleotides for the microarrays. The accession numbers for the viral genomes used are given, along with the gene name and location of the region used for oligonucleotide design.

20	Virus	Gene Name	Genome Location
		Ela	12261542
		Elb_1	32703503
		E2a_2	complement(2408925885)
25	Adenovirus, type 2	E3-1	2760929792
	Accession #J01917	E4 (last exon at 3'-end)	complement(3319332802)
		IX	35764034
		lva2	complement(40815417)
30		DNA Polymerase	complement(51875418)
00		HCMVTRL2 (IRL2)	18932240
		HCMVTRL7 (IRL7)	complement(65956843)
		HCMVUL21	complement(2649727024)
		HCMVUL27	complement(3283134657)
35		HCMVUL33	4325144423
	Cytomegalovirus (CMV)	HCMVUL54	complement(7690380631)
		HCMVUL75	complement(107901110132)
		HCMVUL83	complement(119352121037)
40	Accession #X17403	HCMVUL106	complement(154947155324)
		HCMVUL109	complement(157514157810)
		HCMVUL113	161503162800
		HCMVUL122	complement(169364170599)
		HCMVUL123 (last exon at 3'-end)	complement(171006172225)
45		HCMVUS28	219200220171
		Exon in EBNA-1 RNA	6747767649
	Epstein-Barr virus (EBV)	Exon in EBNA-1 RNA	9836498730
		BRLF1	complement(103366105183)
50		BZLF1 (first of 3 exons)	complement(102655103155)
	Accession # NC_001345	BMLF1	complement(8274384059)
		BALF2	complement(161384164770)
		U16/U17	complement(2625927349)
55		U89	complement(133091135610)
		U90	complement(135664135948)
		U86	complement(125989128136)

(continued)

Virus **Gene Name Genome Location** U83 123528..123821 U22 complement(33739..34347) **Human Herpesvirus 6 (HHV6)** DR2 (DR2L) 791..2653 DR7(DR7L) 5629..6720 Accession #NC_001664 U95 142941..146306 U94 complement(141394..142866) U39 complement(59588..62080) U42 complement(69054..70598) U81 complement(121810..122577) U91 136485..136829

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Table 4: Dependent variables for discovery of gene expression markers of cardiac allograft rejection.

0	Dependent Variable	Description	Number of Rejection Samples	Number of No-Rejection Samples
,	0 vs 1-4 Bx	Grade 0 vs. Grades 1-4, local biopsy reading	65	114
5	s0 vs I B-4 HG	Stable Grade 0 vs Grades 1B-4, highest grade, Grade 1A not included	41	57
	0-1A vs 1B-4 HG	Grades 0 and 1A vs Grades 1B-4, highest grade.	121	58
)	0 vs 3A HG	Grade 0 vs Grade 3A, highest grade. Grades 1A-2 and Grade 3B were not included.	56	29
5	0 vs 1B-4	Grade 0 vs Grades 1B-4, highest grade. Grade 1A was not included.	57	57
	0 vs 1A-4	Grade 0 vs. Grades 1-4,	56	123

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Table 5: Real-time PCR assay chemistries. Various combinations of reporter and quencher dyes are useful for real-time PCR assays.

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Reporter	Quencher
FAM	TAMRA
	BHQ1
TET	TAMRA
	BHQ1
JOE	TAMRA
	BHQ1
HEX	TAMRA
	BHQ1

(continued)

Reporter	Quencher
VIC	TAMRA
	BHQ1
ROX	BHQ2
TAMRA	BHQ2

Table 6: Real-time PCR results for rejection markers

15	
20	
25	
30	
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Gene		Phase 1				Phase 2	2			All Data	l	
Array Probe SEQ IDID	Fold	t-Test	NR	R	Fold	t-Test	NR	R	Fold	t-Test	NR	R
79	1.822	0.01146	6	7	0.63	0.04185	19	15	0.72	0.05632	35	26
3016	1.045	0.41017	12	10					1.001	0.49647	16	15
2766	0.956	0.43918	12	10	0.989	0.48275	19	14	0.978	0.45101	31	24
94	1.601	0.02418	11	10					1.831	0.00094	17	15
36	1.605	0.09781	12	8	2.618	0.01227	18	11	2.808	0.00015	38	23
80	5.395	0.00049	9	6	4.404	0.05464	10	10	2.33	0.02369	29	18
77	1.894	0.01602	10	10	0.537	0.01516	19	15	0.863	0.21987	35	29
2772	1.583	0.06276	10	6	0.714	0.13019	13	10	1.136	0.28841	28	17
102	1.245	0.05079	11	10	1.018	0.42702	17	15	1.117	0.08232	32	28
78	1.324	0.01985	12	9	0.967	0.33851	18	14	1.007	0.46864	38	24
88	1.282	0.14287		7	0.995	0.48504	17	14	1.008	0.47383	30	23
2781					0.492	0.01344	12	12	0.819	0.28555	17	15
101					0.652	0.04317	19	15	0.773	0.09274	29	22
87					4.947	0.02192	18	15	3.857	0.00389	30	23
104	2.292	0.0024	11	8	0.621	0.05152	19	15	0.913	0.34506	30	23
91	1.989	0.07789	11	8	4.047	0.00812	19		3.535	0.00033	37	23
85	0.95	0.43363	12	8	0.699	0.0787	13	13	0.633	0.01486	33	24
2783	0.945	0.48023	10	5	0.852	0.28701	17	10	0.986	0.48609	29	17
2784	2.12	0.00022	12	10	0.498	0.01324	18	13	0.935	0.37356	37	25
3018	1.523	0.1487	12	10	0.84	0.27108	18	13	1.101	0.33276	36	26
3019	1.268	0.21268	6	7	0.981	0.45897	16	10	1.012	0.46612	29	19
2790	0.881	0.17766	11	8	1.22	0.04253	18	10	0.966	0.33826	40	23
3020	1.271	0.10162	12	10	0.853	0.10567	19	13	0.965	0.36499	36	25
2794	1.936	0.00176	13	9	0.717	0.09799	19	14	0.877	0.22295 -	40	25

Table 7: Significance analysis for microarrays for identification of markers of acute rejection. In each case the highest grade from the 3 pathologists was taken for analysis. No rejection and rejection classes are defined. Samples are either used regardless of redundancy with respect to patients or a requirement is made that only one sample is used per patient or per patient per class. The number of samples used in the analysis is given and the lowest FDR achieved is noted.

No Rejection	Rejection	# Samples	Low FDR
All Samples		<u> </u>	
Grade 0	Grade 3A-4	148	1
Grade 0	Grade 1B, 3A-4	158	1.5
Non-redundant with	in class	·	<u>.</u>
Grade 0	Grade 3A-4	86	7
Grade 0	Grade 1B, 3A-4	93	16
Non-redundant (1 sa	ample/patient)	<u> </u>	
Grade 0	Grade 3A-4	73	11

5		ation and r	CD8, CTL effector; channel-forming protein capable of lysing nonspecifically a variety of target cells; clearance of virally infected host cells; and tumor cells;	Induces stromal cells to produce proinflammatory and hematopoietic cytokines; enhances IL6, IL8 and ICAM-1 expression in fibroblasts; osteoclastic bone resorption in RA; expressed in only in activated CD4+T cells	ıatory	Neurogenesis, immune system development, signaling	fense and tivation	Promotes growth of B and T cells	
10	Function	T-cell activation and proliferation	CD8, CTL effector; channel-forming prote capable of lysing nonspecifically a variety o target cells; clearance virally infected host ce and tumor cells;	Induces stromal celliproduce proinflamma and hematopoietic cytokines; enhances IL8 and ICAM-1 expression in fibrobic osteoclastic bone resorption in RA; expressed in only in activated CD4+T celliproduces activated CD4+T celliproduces produced in the structure of the structure o	Proinflammatory cytokine	Neurogenesis, immur system development, signaling	Antiviral defense and immune activation	Promotes g T cells	
15	uc								
	Localization	Secreted	Secreted	Secreted	Secreted	Secreted	Secreted	Secreted	Secreted
20 25	Current UniGene Cluster (Build 156)	Hs.168132	Hs.2200	Hs.41724	Hs.624	Hs.789	Hs.856	Hs.89679	Hs.75415
30	RefSeq Peptide Accession#	NP_000576	NP_005032	NP_002181	NP_ 000575	NP_001502	NP_000610	NP_000577	NP_004039
35	mRNA Accession #	NM_00585	NM_005041	NM_002190	NM_000584	NM_001511	NM_000619	NM_000586	NM_004048
40			forming	vfotoxic sociated 3)		-C motif) la growth ty, alpha)	па		ulin
45	Gene Name	Interleukin 15	Perforin 1 (pore forming protein)	Interleukin 17 (cytotoxic T-lymphocyte-associated serine esterase 8)	Interleukin 8	Chemokine(C-X-C motif) ligand (melanoma growth stimulating activity, alpha)	Interferon, gamma	Interleukin 2	beta 2 microglobulin
50	ne	5	<u>T</u>			CXCL1	JG.		>
55	Array Gene Probe SEQ ID	IL15	79 PRF1	1.17	IL8	Š	IFNG	IL2	B2M

5		Function	Chemoattractant for monocytes, memory T helper cells and eosinophils; causes release of histamine from basophils and activates eosinophils; One of the major HIV-suppressive factors produced by CD8+ cells	Chemotactic factor for CD8+T cells; down-regulates expression of Th1 cytokines, MHC class II Ags, and costimulatory molecules on macrophages; enhances B cell survival, proliferation, and antibody production; blocks NF kappa B, JAK-STAT regulation;	TH2, cytokine, stimulates CTL	Proliferation of lymphoid progenitors
15		Localization	Secreted	Secreted	Secreted	Secreted
20		Current UniGene Cluster (Build 156)	Hs.241392	Hs.193717	Hs.73917	Hs.72927
30	(continued)	RefSeq Peptide Accession#	NP_002976	NP_000563	085000_AN	NP_000871
35)	mRNA Accession #	NM_002985	NM_000572	000589 NM_000589	NM_000880
40			motif) ligand			
45		Gene Name	Chemokine (C-C motif) ligand 5 (RANTES, SCYAS)	Interleukin 10	Interleukin 4	Interleukin 7
50		Gene	CCL5	0		
55		Array Ge Probe SEQ ID	3	110	80 11.4	IL7 IL7

55	50	40	35	30 interest	20 25	15	5
Array Probe SEQ ID	Gene	Gene Name	mRNA Accession#	RefSeq Peptide Accession #	Current UniGene Cluster (Build 156)	Localization	Function
	CXCL10	Chemokine (C-X-C motif) ligand 10, SCYB10	NM_001565	NP_001556	Hs.2248	Secreted	Stimulation of monocytes; NK and T cell migration, modulation of adhesion molecule expression
	CCL17	Chemokine (C-C motif) ligand 17	NM_002987	NP_002978	Hs.66742	Secreted	T cell development, trafficking and activation
101	KLRF1	Killer cell lectin-like receptor subfamily F, member I	NM_016523	NP_057607	Hs.183125	Secreted	Induction of IgE, IgG4, CD23, CD72, surface IgM, and class II MHC antigen in B cells
	971	Interleukin 6	009000 ⁻ MN	NP_ 000591	Hs.93913	Secreted	B cell maturation
	CCL4	Chemokine (C-C motif) ligand 4	NM_002984 4	NP_ 002975	Hs.75703	Secreted	Inflammatory and chemokinetic properties; one of the major HIV-suppressive factors produced by CD8+ T cells
	GZMB	Granzyme B (granzyme 2, cytotoxic T-lymphocyte-associated serine esterase I)	NM_004131	NP_004122	Hs.1051	Secreted	Apoptosis; CD8, CTL effector
	OID_4789	KIAA0963 protein	NM_014963	NP_055778	Hs.7724	Secreted	Proinflammatory; chemoattraction and activation of neutrophits
	XCL1	Chemokine (C motif) ligand 1 (SCYC2)	NM_002995	NP_002986	Hs.3195	Secreted	Chemotactic factor for lymphocytes but not monocytes or neutrophils

5	Function	Transcription factor; promotes B cell maturation, represses human beta-1FN gene expression	TH1 differentiation, transcription factor	Folate metabolism	T cell mediated immune response	CD8, CTL effector proapoptotic	CTL mediated killing	Angiogenesis, cell migration, synthesis of inflammatory prostaglandins	Transports antigens into ER for association with MHC class 1 molecules	Antibody subunit
15	Localization	Nuclear	Nuclear	Mitochondrial	Membrane-bound and soluble forms	Membrane-bound and soluble forms	Membrane-bound and soluble forms	Membrane-associated	ER membrane	Cytoplasmic and secreted forms
20	Current UniGene Cluster (Build 156)	Hs.388346	Hs.272409	Hs.154672	Hs.1724	Hs.2007	Hs.2299	Hs.196384	Hs.352018	Hs.300697
30	(continued) RefSeq Peptide Accession#	NP_001189	NP_037483	NP_006627	NP_000408	NP_000630	NP00_4922	NP_000954	NP_000584	
35	mRNA Accession #	NM_001198 with	NM_013351	NM_006636	NM_000417	NM_000639 (ligand)	NM_004931 1	NM_000963	NM_000593	BC032249
40		taining 1, with		hydrofolate (NAD+ thenyltetrahydr drolase	ceptor, alpha	factor (ligand) mber 6	sta polypeptide	ndoperoxide staglandin G/H /clooxygenase)	ATP-binding mily B	n heavy
45	Gene Name	PR domain containing 1, with ZNF domain	T-box 21	Methylene tetrahydrofolate dehydrogenase (NAD+ dependent), methenyltetrahydr ofolate cyclohydrolase	Interleukin 2 receptor, alpha	Tumor necrosis factor (ligand) superfamily, member 6	CD8 antigen, beta polypeptide 1 (p37)	Prostaglandin-endoperoxide synthasc 2 (prostaglandin G/H synthase and cyclooxygenase)	Transporter 1, ATP-binding cassette, subfamily B (MDR1/TAP)	Immunoglobulin heavy constant mu
50	Gene	PRDM1	TBX21	МТНБD2	IL2RA	TNFSF6	CD8B1	PTGS2	TAP1	МНЭ
55	Array G Probe SEQ ID	ш.	2781 T	88	=	T 77		ш	Т	

5	Function	Glycolysis and gluconeogenesis (cytoplasmic); neurotrophic factor (secreted)	Controls actin filament assembly/disassembly	Mediator of stress-activated signals; Serine/Thr Kinase, activated p38	Processing of MHC class I antigens	Signal transduction	Intracellularkinase, T-cell proliferation and differentiation	Nucleocytoplasmic transport	CD8 T activation, signal transduction	B-cell proliferation, IgE production, immunoglobulin class switching; expressed on CD4+ and CD8+ T cells
15	Localization	Cytoplasmic and secreted forms	Cytoplasmic and secreted forms	Cytoplasmic and nuclear	Cytoplasmic	Cytoplasmic	Cytoplasmic	Cytoplasmic	Cytoplasmic	Cellular membrane
25	Current UniGene Cluster(Build 156)	Hs.409162	Hs.290070	Hs.199263	Hs.180062,	Hs.49587 .	Hs.211576	Hs.301553	Hs. 103527	Hs.652
% (continued)	RefSeq Peptide Accession#	NP_000166	NP_000168	NP_037365	Hs.180062	NP_004802			NP_003966	NP_000065
35	mRNA Accession #	NM_000175	NM_000177	NM_013233	AK092738	NM_004811	L10717	AW021037	NM_003975	NM_000074
40	Gene Name	Glucose phosphate NM_000175 isomerase	Gelsolin (amyloidosis, Finnish type)	Serine threonine kinase 39 (STE20/SPS1 homolog, yeast)	Proteasome (prosome, macropain) subunit, beta type, 8 (large multifunctional protease 7)	axin	IL2-inducible T-cell kinase	Karyopherin alpha 6 (importin alpha 7)	SH2 domain protein	Tumor necrosis factor (ligand) superfamily, member 5 (hyper-IgM syndrome)
50	Gene	Glucc NM_C	Gelsc type)	Serin (STE)		Leupaxin	IL2-in			
55	Array Gene Probe SEQ ID	GP1	2783 GSN	STK39	PSMB8	NXAT	ПК	90 KPNA6	2794 SH2D2A	TNF8F5

5		Function	Activation of lymphocytes, monocytes, and platelets	Signalling component of many interleukin receptors (IL2,IL4,IL7,IL9, and IL15),	B-cell lymphopoiesis, leukocyte migration, angiogenesis; mediates intracellular calcium flux	Signal transduction; B lymphocyte development, activation, and differentiation	Cell-cell and cell-matrix interactions	Antigen recognition	Negative regulation ofT cell activation, expressed by activated T cells	CD8 T-cell specific marker and class I MHC receptor	Antigen presentation
15		Localization	Cellular membrane	Cellular membrane	Cellular membrane	Cellular membrane	Cellular membrane	Cellular membrane	Cellular membrane	Cellular membrane	Cellular membrane
2025		Current UniGene Cluster(Build 156)	Hs.82401	Hs.84	Hs.89414	Hs.96023		Hs.300697	Hs.247824	Hs.85258	Hs.308026
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40			(p60, early gen)	eceptor, gamma ined sncy)	-X-C motif)		ntegrin, beta I (fibronectin receptor, beta polypeptide, antigen CD29 includes MDF2, MSK12)	T cell receptor beta, constant region	mphocyte- xtein 4	CD8 antigen, alpha polypeptide (p32)	npatibility s II, DR beta 1
45		Gene Name	CD69 antigen (p60, early activation antigen)	Interleukin 2 receptor, gamr (severe combined immunodeficiency)	Chemokine (C-X-C motif) receptor 4	CD19 antigen	ntegrin, beta I (fibronectin receptor, beta polypeptide, antigen CD29 includes MD MSK12)	T cell receptor region	Cytotoxic T-lymphocyte- associated protein 4	CD8antigen, a (p32)	Major histocompatibility complex, class II, DR beta
50		Gene	CD69	IL2RG	CXCR4	CD19	4TGBI	TRB	CTLA4	CD8A	HLA-DRB1
55		Array Probe SEQ ID				2766					

5		Function	T-cell marker, couples antigen recognition to several intracellular signal-transduction pathways	Cell adhesion and recognition	All leukocytes; cell-cell adhesion, signaling	T cell activation	B cell proliferation	NK cells marker	B-cell activation, plasma cell development		T cell activation, signal transduction, T-B cell adhesion
15		Localization	Cellular membrane	Cellular membrane	Cellular membrane	Cellular membrane	Cellular membrane	Cellular membrane	Cellular membrane	Cellular membrane	Cellular membrane
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40			zeta r3 complex)		(antigen CD nphocyte ated antigen 1; de)	regulator 1, nsporting, otein a isoform		n chromosome expressed	nning family A, 20	amma constant	55)
45		Gene Name	CD3Z antigen, zeta polypeptide (TiT3 complex)	Actin, beta	Integrin, alpha L (antigen CD 11A (p1 80), lymphocyte function-associated antigen 1 alpha polypeptide)	T-cell, immune regulator 1, ATPasc, H+ transporting, lysosomal V0 protein a isoform 3	CD72 antigen	DNA segment on chromosome 1 (unique) 2489 expressed sequence	Membrane-spanning 4-domains, subfamily A, member 1, CD20	T cell receptor gamma constant 2	CD4 antigen (p55)
50		Gene	CD3Z	ACTB	ITGAL	TCIRG1	CD72	D1252489E	MS4A1	TCRGC2	CD4
55		Array Probe SEQ ID	2772			78					

5		Function	Integrin activation, cytoskeletal changes and chemotactic migration of leukocytes	Cell adhesion; receptor that inhibits the proliferation of normal and leukemic myeloid cells	Cell adhesion, membrane transport, signaling transduction, permeability	NK cells and CTLs, costim with MHC I	NK cell regulation	Antigen presentation	T cell activation
15		Localization	Cellular membrane	Cellular membrane	Cellular membrane	Cellular membrane	Cellular membrane	Cellular membrane	Cellular membrane
20		Current UniGene Cluster (Build 156)	Hs. 198252	Hs.83731	Hs.82685	Hs.81743	Hs.41682	Hs.377850	Hs.155975
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40 45		Gene Name	Chemokine(C-X-C motif) receptor 3, GPR9	CD33 antigen (gp67)	CD47 antigen (Rh-related antigen, integin-associated signal transducer)	Natural killer cell receptor, immunoglobulin superfamily member	Killer cell lectin-like receptor subfamily D, member I	Major histocompatibility complex, class 1, F	Protein tyrosine phosphatase, receptor type, C-associated protein
50		Gene Gene	CXCR3 Chem recept	CD33 CD33	CD47 CD47 antige signal	BY55 Natural P immunog	KLRD1 Killer subfar	HLA-F Major compl	PTPRCAP Protein recepto protein
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Ala	Leu	Ala	Met 420	val	Leu	Аlа	Glu	Asn 425	Arg	Pro	Ala	val	Lys 430	Ala	Ala
Gln	Asn	Glu 435	Ile	Trp	Gln	Thr	Phe 440	Phe	Arg	Gly	Arg	Tyr 445	Leu	Leu	Leu
Leu	Met 450	Gly	Leu	Phe	Ser	Ile 455	Туг	Thr	Gly	Phe	11e 460	туг	Asn	Glu	Cys
Phe 465	Ser	Arg	Ala	Thr	Ser 470	Ile	Phe	Pro	Ser	Gly 475	Trp	Ser	val	Ala	Ala 480
Met	Ala	Asn	Gln	Ser 485	Gly	Trp	Ser	Asp	Ala 490	Phe	Leu	Ala	Gln	ніs 495	Thr
Met	Leu	Thr	Leu	Asp	Pro	Asn	val	Thr	Gly	۷a۱	Phe	Leu	Gly	Pro	Tyr

500 505 510

Pro Phe Gly Ile Asp Pro Ile Trp Ser Leu Ala Ala Asn His Leu Ser Phe Leu Asn Ser Phe Lys Met Lys Met Ser Val Ile Leu Gly Val Val 530 540 His Met Ala Phe Gly Val Val Leu Gly Val Phe Asn His Val His Phe 545 550 560 10 Gly Gln Arg His Arg Leu Leu Glu Thr Leu Pro Glu Leu Thr Phe 565 570 575 Leu Leu Gly Leu Phe Gly Tyr Leu Val Phe Leu Val Ile Tyr Lys Trp 580 585 590 15 Leu Cys Val Trp Ala Ala Arg Ala Ala Ser Ala Pro Ser Ile Leu Ile 595 600 605 His Phe Ile Asn Met Phe Leu Phe Ser His Ser Pro Ser Asn Arg Leu 610 615 620 20 Leu Tyr Pro Arg Gln Glu Val Val Gln Ala Thr Leu Val Val Leu Ala 625 630 635 640 Leu Ala Met Val Pro Ile Leu Leu Leu Gly Thr Pro Leu His Leu Leu 645 650 655 His Arg His Arg Arg Arg Leu Arg Arg Arg Pro Ala Asp Arg Gln Glu 660 670 Glu Asn Lys Ala Gly Leu Leu Asp Leu Pro Asp Ala Ser Val Asn Gly $675 \hspace{1.5cm} 680 \hspace{1.5cm} 685$ 30 Trp Ser Ser Asp Glu Glu Lys Ala Gly Gly Leu Asp Asp Glu Glu Glu 690 700 Ala Glu Leu Val Pro Ser Glu Val Leu Met His Gln Ala Ile His Thr 705 710 715 720 35 Ile Glu Phe Cys Leu Gly Cys Val Ser Asn Thr Ala Ser Tyr Leu Arg Leu Trp Ala Leu Ser Leu Ala His Ala Gln Leu Ser Glu Val Leu Trp 740 745 750 40 Ala Met Val Met Arg Ile Gly Leu Gly Leu Gly Arg Glu Val Gly Val 755 760 765 45 Ala Ala Val Val Leu Val Pro Ile Phe Ala Ala Phe Ala Val Met Thr $770 \hspace{1.5cm} 780$ Val Ala Ile Leu Leu Val Met Glu Gly Leu Ser Ala Phe Leu His Ala 785 790 795 800 50 Leu Arg Leu His Trp Val Glu Phe Gln Asn Lys Phe Tyr Ser Gly Thr 805 810 815 Gly Tyr Lys Leu Ser Pro Phe Thr Phe Ala Ala Thr Asp Asp 820 825 830

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Gly	Leu	٧a٦	Asp 340	Tyr	Thr	Leu	Glu	Рго 345	Leu	His	۷al	Leu	Leu 350	Asp	Ser
Gln	Asp	Pro 355	Arg	Arg	Glu	Ala	Leu 360	Arg	Arg	Ala	Leu	Ser 365	Gln	Туг	Leu
Thr	Asp 370	Arg	Ala	Arg	тгр	Arg 375	Asp	Cys	Ser	Arg	Pro 380	Cys	Pro	Pro	Gly
Arg 385	Gln	Lys	Ser	Pro	Arg 390	Asp	Pro	Cys	Gln	Cys 395	val	Cys	His	G1y	Ser 400
Ala	val	Thr	Thr	G1n 405	Asp	Cys	Cys	Pro	Arg 410	Gln	Arg	Gly	Leu	Ala 415	Gln
Leu	G1u	val	Thr 420	Phe	Ile	Gln	Ala	Trp 425	Ser	Leu	Trp	Gly	Asp 430	Тгр	Phe
Thr	Ala	Thr 435	Asp	Ala	Туг	٧al	Lys 440	Leu	Phe	Phe	Gly	G1y 445	Gln	Glu	Leu
Arg	Thr 450	Ser	Thr	∨al	Тгр	Asp 455	Asn	Asn	Asn	Pro	11e 460	Тгр	Ser	val	Arg
Leu 465	Asp	Phe	Gly	Asp	Va1 470	Leu	Leu	Ala	Thr	G1y 475	Gly	Pro	Leu	Arg	Leu 480
Gln	val	Trp	Asp	G]n 485	Asp	Ser	Gly	Arg	Asp 490	Asp	Asp	Leu	Leu	Gly 495	Thr
Cys	Asp	Gln	Ala 500	Pro	Lys	Ser	Gly	Ser 505	His	Glu	val	Arg	Cys 510	Asn	Leu
Asn	нis	Gly 515	His	Leu	Lys	Phe	Arg 520	Tyr	His	Ala	Arg	Cys 525	Leu	Pro	His
Leu	Gly 530	Gly	Gly	Thr	Cys	Leu 535	Asp	Tyr	val	Pro	G1n 540	Met	Leu	Leu	Gly
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<213> Homo sapiens

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		Glu	Ile	Ile 35	Lys	Thr	Leu	Asn	Ser 40	Leu	Thr	Glu	Gln	Lys 45	Thr	Leu	Cys
10		Thr	G1u 50	Leu	Thr	٧al	Thr	Asp 55	Ile	Phe	Ala	Ala	Ser 60	Lys	Asn	Thr	Thr
		Glu 65	Lys	Glu	Thr	Phe	Cys 70	Arg	Ala	Ala	Thr	Va1 75	Leu	Arg	Gln	Phe	Tyr 80
15		Ser	His	His	G1u	Lys	Asp	Thr	Arg	Cys	Leu	Gly	Ala	Thr	Ala	Gln	Gln
						85					90					95	
20		Phe	His	Arg	His 100	Lys	Gln	Leu	Ile	Arg 105	Phe	Leu	Lys	Arg	Leu 110	ASP	Arg
		Asn	Leu	Trp 115	Gly	Leu	Ala	Gly	Leu 120	Asn	Ser	Cys	Pro	Val 125	Lys	Glu	Ala
25		Asn	Gln 130	Ser	Thr	Leu	Glu	Asn 135	Phe	Leu	Glu	Arg	Leu 140	Lys	Thr	Ile	Met
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		Leu	val	va1 35	туг	Pro	Тгр	Thr	G]n 40	Arg	Phe	Phe	Glu	Ser 45	Phe	Gly	Asp
10		Leu	ser 50	Thr	Pro	Asp	Ala	va1 55	Met	Glу	Asn	Pro	Lys 60	٧a٦	Lys	Ala	His
		Gly 65	Lys	Lys	val	Leu	G]y 70	Ala	Phe	Ser	Asp	G]y 75	Leu	Ala	His	Leu	Asp 80
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		Leu	His	val	Asp 100	Pro	Glu	Asn	Phe	Arg 105	Leu	Leu	Gly	Asn	Val 110	Leu	Val
20		Cys	val	Leu 115	Ala	His	His	Phe	Gly 120	Lys	Glu	Phe	Thr	Pro 125	Pro	val	Gln
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		Glu	Gly	Met 35	Glu	Glu	Ala	Arg	Asn 40	Cys	Gly	Lys	Gln	Leu 45	Lys	Ala	Leu
10		Asn	Phe 50	Glu	Phe	Asp	Leu	va1 55	Phe	Thr	Ser	val	Leu 60	Asn	Arg	Ser	IJe
		His 65	Thr	Ala	Тгр	Leu	11e 70	Leu	Glu	Glu	Leu	Gly 75	Gln	Glu	Trp	val	Pro 80
15		۷al	Glu	Ser	Ser	Trp 85	Arg	Leu	Asn	Glu	Arg 90	His	Туг	Gly	Ala	Leu 95	Ile
		Gly	Leu	Asn	Arg 100	Glu	Gln	Met	Ala	Leu 105	Asn	His	Gly	Glu	Glu 110	Gln	val
20		Arg	Leu	Trp 115	Arg	Arg	Ser	туг	Asn 120	val	Thr	Pro	Pro	Pro 125	Ile	Glu	Glu
		Ser	Нis 130	Pro	туг	Tyr	Gln	Glu 135	Ile	Туг	Asn	Asp	Arg 140	Arg	Туг	Lys	val
25		Cys 145	Asp	٧a٦	Pro	Leu	Asp 150	Gln	Leu	Pro	Arg	Ser 155	Glu	Ser	Leu	Lys	Asp 160
20		∨al	Leu	Glu	Arg	Leu 165	Leu	Pro	Туг	Тгр	Asn 170	Glu	Arg	Ile	Ala	Pro 175	Glu
30		val	Leu	Arg	Gly 180	Lys	Thr	Ile	Leu	Ile 185	Ser	Ala	His	Gly	Asn 190	Ser	Ser
35		Arg	Ala	Leu 195	Leu	Lys	His	Leu	G]u 200	Gly	Ile	Ser	Asp	G1u 205	Asp	Ile	Ile
		Asn	Ile 210	Thr	Leu	Pro	Thr	Gly 215	val	Pro	Ile	Leu	Leu 220	Glu	Leu	Asp	Glu
40		Asn 225	Leu	Arg	Ala	∨al	Gly 230	Pro	His	Gln	Phe	Leu 235	Gly	Asp	Gln	Glu	Ala 240
		Ile	Gln	Ala	Ala	11e 245	Lys	Lys	val	Glu	Asp 250	Gln	Gly	Lys	val	Lys 255	Gln
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	val	Ile	Ser 35	Gly	Arg	Lys	Leu	Ala 40	Gln	Gln	Ile	Lys	G]n 45	Glu	۷al	Arg
10	Gln	Glu 50	val	Glu	Glu	Тгр	Va1 55	Ala	Ser	Gly	Asn	Lys 60	Arg	Pro	His	Leu
	Ser 65	۷al	Ile	Leu	٧a٦	Gly 70	Glu	Asn	Pro	Ala	Ser 75	His	Ser	Туг	val	Leu 80
15	Asn	Lys	Thr	Arg	Ala 85	Ala	Ala	val	val	G] y 90	Ile	Asn	Ser	Glu	T h r 95	Ile
	Met	Lys	Pro	Ala 100	ser	Ile	Ser	Glu	Glu 105	Glu	Leu	Leu	Asn	Leu 110	Ile	Asn
20	Lys	Leu	Asn 115	Asn	Asp	Asp	Asn	val 120	Asp	Gly	Leu	Leu	va1 125	Gln	Leu	Pro
	Leu	Pro 130	Glu	His	Ile	Asp	Glu 135	Arg	Arg	Ile	Cys	Asn 140	Ala	val	Ser	Pro
25	Asp 145	Lys	Asp	val	Asp	Gly 150	Phe	His	val	Ile	Asn 155	Val	Gly	Arg	Met	Cys 160
	Leu	Asp	Gln	Туг	Ser 165	Met	Leu	Pro	Ala	Thr 170	Pro	Trp	Gly	Val	Trp 175	Glu
30	Ile	Ile	Lys	Arg 180	Thr	Gly	Ile	Pro	Thr 185	Leu	Gly	Lys	Asn	val 190	val	۷al
05	Ala	Gly	Arg 195	Ser	Lys	Asn	val	Gly 200	Met	Pro	Ile	Ala	Met 205	Leu	Leu	His
35	Thr	Asp 210	Gly	Ala	нis	Glu	Arg 215	Pro	Gly	Gly	Asp	Ala 220	Thr	val	Thr	ΙΊe
40	Ser 225	ніѕ	Arg	Туг	Thr	Pro 230	Lys	Glu	Gln	Leu	Lys 235	Lys	His	Thr	Ile	Leu 240
	Ala	Asp	Ile	val	Ile 245	Ser	Ala	Ala	Gly	Ile 250	Pro	Asn	Leu	Ile	Thr 255	Ala
45	Asp	Met	Ile	Lys 260	Glu	Gly	Ala	Ala	va1 265	Ile	Asp	val	Glу	Ile 270	Asn	Arg
	val	His	Asp 275	Pro	val	Thr	Ala	Lys 280	Pro	Lys	Leu	val	G]y 285	Asp	va1	Asp
50	Phe	G1u 290	G1y	val	Arg	Gln	Lys 295	Ala	Gly	туг	Ile	Thr 300	Pro	val	Pro	Gly
	G]y 305	٧al	Gly	Pro	Met	Thr 310	۷al	Ala	мet	Leu	Met 315	Lys	Asn	Thr	Ile	11e 320
55	Ala	Αla	Lys	Lys	va1 325	Leu	Arg	Leu	Glu	G] u 330	Arg	Glu	val	Leu	Lys 335	ser
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Gly	Ala	Gly	Ser 20	ASP	Ala	Gln	Gly	Pro 25	Gln	Phe	Gly	Тгр	Asp 30	His	Ser
Leu	His	Lys 35	Arg	Lys	Arg	Leu	Pro 40	Pro	val	Lys	Arg	Ser 45	Leu	val	Туг
Tyr	Leu 50	Lys	Asn	Arg	Glu	va1 55	Arg	Leu	Gln	Asn	Glu 60	Thr	Ser	туг	Ser
Arg 65	Va1	Leu	His	Gly	Туг 70	Ala	Ala	Gln	Gln	Leu 75	Pro	Ser	Leu	Leu	Lys 80
Glu	Arg	Glu	Phe	His 85	Leu	Gly	Thr	Leu	Asn 90	Lys	val	Phe	Ala	Ser 95	Gln
тгр	Leu	Asn	Нis 100	Arg	Gln	val	val	Cys 105	Gly	Thr	Lys	Cys	Asn 110	Thr	Leu
Phe	val	Val 115	Asp	val	Gln	Thr	Ser 120	G]n	Ile	Thr	Lys	Ile 125	Pro	Ile	Leu
Lys	Asp 130	Arg	Glu	Pro	Gly	Gly 135	val	Thr	Gln	Gln	Gly 140	Cys	Gly	Ile	His
Ala 145	Ile	Glu	Leu	Asn	Pro 150	Ser	Arg	Thr	Leu	Leu 155	Ala	Thr	Gly	Gly	Asp 160
Asn	Pro	Asn	Ser	Leu 165	Ala	Ile	Tyr	Arg	Leu 170	Pro	Thr	Leu	Asp	Pro 175	val
Cys	val	Gly	Asp 180	Asp	Gly	His	Lys	Asp 185	Trp	Ile	Phe	Ser	Ile 190	Ala	Тгр
Ile	Ser	Asp 195	Thr	Met	Ala	val	Ser 200	Gly	Ser	Arg	Asp	G]y 205	Ser	Met	Gly
Leu	Trp 210	Glu	val	Thr	Asp	Asp 215	val	Leu	Thr	Lys	Ser 220	Asp	Ala	Arg	ніѕ
Asn 225	val	Ser	Arg	۷al	Pro 230	val	Туг	Ala	His	Ile 235	Thr	His	Lys	Ala	Leu 240
Lys	Asp	Ile	Pro	Lys 245	Glu	Asp	Thr	Asn	Pro 250	Asp	Asn	Cys	Lys	Va1 255	Arg
Ala	Leu	Ala	Phe 260	Asn	Asn	Lys	Asn	Lys 265	Glu	Leu	Gly	Ala	va1 270	Ser	Leu
Asp	Gly	Tyr 275	Phe	His	Leu	Trp	Lys 280	Ala	Glu	Asn	Thr	Leu 285	Ser	Lys	Leu
Leu	Ser 290	Thr	Lys	Leu	Pro	Tyr 295	Cys	Arg	Glu	Asn	va1 300	Cys	Leu	Ala	туг
G]y 305	Ser	Glu	Trp	Ser	Val 310	Tyr	Ala	Val	Gly	Ser 315	Gln	Ala	нis	۷al	Ser 320
Phe	Leu	Asp	Pro	Arg 325	G]n	Pro	Ser	туг	Asn 330	val	Lys	Ser	val	Cys 335	Ser
Arg	Glu	Arg	Gly 340	Ser	Gly	Ile	Arg	Ser 345	val	Ser	Phe	Туг	Glu 350	His	Ile

		Ile	Thr	va1 355	Gly	Thr	Gly	Gln	Gly 360	Ser	Leu	Leu	Phe	Tyr 365	Asp	Ile	Arg
5		Ala	G1n 370	Arg	Phe	Leu	Glu	G1u 375	Arg	Leu	Ser	Ala	Cys 380	туг	Gly	Ser	Lys
		Pro 385	Arg	Leu	Ala	Gly	Glu 390	Asn	Leu	Lys	Leu	Thr 395	Thr	Gly	Lys	Gly	Trp 400
10		Leu	Asn	His	Asp	G1u 405	Thr	Тгр	Arg	Asn	туг 410	Phe	Ser	Asp	ıle	ASP 415	Phe
		Phe	Pro	Asn	Ala 420	val	туг	Thr	His	Cys 425	Tyr	Asp	Ser	Ser	Gly 430	Thr	Lys
15		Leu	Phe	Val 435	Ala	Gly	Gly	Pro	Leu 440	Pro	Ser	Gly	Leu	ніs 445	Gly	Asn	Tyr
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		Thr	Leu	His 35	Trp	Туг	Lys	Ile	Leu 40	Leu	Gly	Ile	Ser	Gly 45	Thr	val	Asn
35		Gly	11e 50	Leu	Thr	Leu	Thr	Leu 55	Ile	Ser	Leu	Ile	Leu 60	Leu	val	ser	Gln
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5		Туг	Glu	Asp	Thr	Gly 85	Asp	Leu	Lys	val	Asn 90	Asn	Gly	Thr	Arg	Arg 95	Asn
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		Ser	Asn 130	Glu	Met	Lys	Ser	Trp 135	Ser	Asp	Ser	Туг	Val 140	туг	Cys	Leu	Glu
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		Asp	ser	Lys 195	Ile	Phe	Phe	Ile	Lys 200	Gly	Pro	Ala	Lys	G1u 205	Asn	Ser	Cys
25		Ala	Ala 210	Ile	Lys	Glu	Ser	Lys 215	Ile	Phe	Ser	Glu	Thr 220	Cys	Ser	Ser	Val
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	val	Arg	Asp 35	Pro	Thr	Ser	Asn	Lys 40	Gln	Gln	Arg	Pro	Va1 45	Pro	Glu	Ser
10	Gln	Leu 50	Leu	Pro	Gly	Gln	Arg 55	Phe	Gln	Thr	Lys	Asp 60	Pro	Glu	Glu	Gln
	Gly 65	Asp	Ile	va1	۷al	Ala 70	Leu	Туг	Pro	Туг	Asp 75	Gly	Ile	нis	Pro	Asp 80
15	Asp	Leu	Ser	Phe	Lys 85	Lys	Gly	Glu	Lys	Met 90	Lys	۷al	Leu	Glu	Glu 95	His
	Gly	Glu	Trp	Trp 100	Lys	Ala	Lys	Ser	Leu 105	Leu	Thr	Lys	Lys	Glu 110	Gly	Phe
20	Ile	Pro	Ser 115	Asn	Tyr	val	Ala	Lys 120	Leu	Asn	Thr	Leu	Glu 125	Thr	Glu	Glu
	Trp	Phe	Phe	Lys	Asp	Ile	Thr	Arg	Lys	Asp	Ala	Glu	Arg	G1n	Leu	Leu
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		130					135					140				
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10	Gly	Asp	٧a٦	Ile 180	Lys	His	Туг	Lys	Ile 185	Arg	Ser	Leu	Asp	Asn 190	Gly	Gly
	Туг	Туг	Ile 195	Ser	Pro	Arg	Ile	Thr 200	Phe	Pro	Cys	Ile	Ser 205	Asp	Met	Ile
15	Lys	His 210	туг	Gln	Lys	Gln	Ala 215	Asp	Gly	Leu	Cys	Arg 220	Arg	Leu	Glu	Lys
	Ala 225	Cys	Ile	Ser	Pro	Lys 230	Pro	Gln	Lys	Pro	Trp 235	Asp	Lys	Asp	Ala	Trp 240
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	Gln	Phe	Gly	G]u 260	val	Тгр	Met	Gly	Tyr 265	туг	Asn	Asn	Ser	Thr 270	Lys	val
25	Ala	val	Lys 275	Thr	Leu	Lys	Pro	Gly 280	Thr	Met	Ser	۷al	G]n 285	Ala	Phe	Leu
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Claims

- 1. Use of one or more genes to assess cardiac allograft rejection in an individual, wherein the expression level of said one or more genes is detected in a blood sample from said patient to assess cardiac allograft rejection versus non-rejection and wherein said one or more genes comprise(s) a nucleotide sequence selected from the group consisting of SEQ ID NO: 86, SEQ ID NO: 87, SEQ ID NO: 94, and SEQ ID NO: 107 and wherein said expression level is detected by measuring the RNA level expressed by said nucleic acid.
- 2. The use according to claim 1, wherein the expression level is detected in peripheral blood leukocytes.

- The use according to any one of claims 1-2, wherein said one or more genes comprise(s) the nucleotide sequence SEQ ID NO: 86.
- The use according to any one of claims 1-2, wherein said one or more genes comprise(s) the nucleotide sequence 30 SEQ ID NO: 87.
 - 5. The use according to any one of claims 1-2, wherein said one or more genes comprise(s) the nucleotide sequence SEQ ID NO: 94.
- 35 6. The use according to any one of claims 1-2, wherein said one or more genes comprise(s) the nucleotide sequence SEQ ID NO: 107.
 - 7. The use according to any one of claims 1-2, wherein said one or more genes comprise(s) the nucleotide sequences SEQ ID NO: 86 and SEQ ID NO: 87.

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- 8. The use according to any one of claims 1-2, wherein the genes comprise the nucleotide sequences SEQ ID NO: 86, SEQ ID NO: 87, SEQ ID NO: 94, and SEQ ID NO: 107.
- The use according to any one of claims 1-8, wherein said RNA is isolated from a sample prior to detecting said RNA 45 level expressed by said gene.
 - 10. The use according any one of claims 1-9, wherein said RNA level is detected by PCR.
 - 11. The use according any one of claims 1-9, wherein said RNA level is detected by hybridization.

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- 12. The use according to claim 11, wherein said RNA level is detected by hybridization to an oligonucleotide.
- 13. The use according to claim 12, wherein said oligonucleotide comprises DNA, RNA, cDNA, PNA, genomic DNA, or synthetic oligonucleotides.

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Patentansprüche

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- 1. Verwendung eines oder mehrerer Gene zur Feststellung einer Herz-Allograftabstoßung in einem Patienten, wobei der Expressionsgrad des einen oder der mehreren Gene in einer Blutprobe des Patienten ermittelt wird, um die Herz-Allograftabstoßung mit der Nichtabstoßung zu vergleichen, wobei das mindestens eine Gen eine Nukleotidsequenz umfasst, die ausgewählt ist aus der Gruppe, welche aus SEQ ID NR.: 86, SEQ ID NR.: 87, SEQ ID NR.: 94 und SEQ ID NR.: 107 besteht und wobei der Expressionsgrad durch Messen der Menge der von der Nukleinsäure exprimierten RNA festgestellt wird.
- 10 2. Verwendung nach Anspruch 1, wobei der Expressionsgrad in Leukozyten des peripheren Bluts ermittelt wird.
 - 3. Verwendung nach einem der Ansprüche 1-2, wobei das mindestens eine Gen die Nukleotidsequenz SEQ ID NR.: 86 umfasst.
- 4. Verwendung nach einem der Ansprüche 1-2, wobei das mindestens eine Gen die Nukleotidsequenz SEQ ID NR.: 87 umfasst.
 - Verwindung nach einem der Ansprüche 1-2, wobei das mindestens eine Gen die Nukleotidsequenz SEQ ID NR.: 94 umfasst.
 - **6.** Verwendung nach einem der Ansprüche 1-2, wobei das mindestens eine Gen die Nukleotidsequenz SEQ ID NR.: 107 umfasst.
- 7. Verwindung nach einem der Anspruche 1-2, wobei das mindestens eine Gen die Nukleotidsequenzen SEQ ID NR.:86 und SEQ ID NR.: 87 umfasst.
 - **8.** Verwindung nach einem der Anspruche 1-2, wobei das mindestens eine Gen die Nukleotidsequenzen SEQ ID NR.: 86, SEQ ID NR.: 94 und SEQ ID NR.: 107 umfasst.
- **9.** Verwerdung nach einem der Anspruche 1-8, wobei die RNA aus einer Probe isoliert wird, bevor die Menge der von dem Gen exprimierten RNA ermittelt wird.
 - 10. Verwerdung nach einem der Ansprüche 1-9, wobei die RNA-Menge mittels PCR ermittelt wird.
- 11. Verwendung nach einem der Ansprüche 1-9, wobei die RNA-Menge durch Hybridisierung ermittelt wird.
 - 12. Verwerdung nach Anspruch 11, wobei die RNA-Menge durch Hybridisierung mit einem Oligonucleotid ermittelt wird.
- **13.** Verwendung nach Anspruch 12, wobei das Oligonucleotid DNA, RNA, cDNA, PNA, genomische DNA oder synthetische Oligonukleotide umfasst.

Revendications

- 45 1. Utilisation d'un ou plusieurs gènes pour l'évaluation du rejet d'une allogreffe cardiaque chez un individu, dans laquelle le niveau d'expression du ou desdits gènes est détecté dans un échantillon sanguin dudit patient afin d'évaluer le rejet d'une allogreffe cardiaque par rapport à son non-rejet, et dans laquelle le ou lesdits gènes comprennent une séquence de nucléotides choisie dans le groupe comprenant les séquences SEQ ID n° 86, SEQ ID n° 87, SEQ ID n° 94 et SEQ ID n° 107 et dans laquelle ledit niveau d'expression est détecté en mesurant le niveau d'ARN exprimé par ledit acide nucléique.
 - 2. Utilisation selon la revendication 1, dans laquelle le niveau d'expression est détecté dans des leucocytes de sang périphérique.
- ⁵⁵ **3.** Utilisation selon l'une quelconque des revendications 1 à 2, dans laquelle le ou lesdits gènes comprennent la séquence de nucléotides SEQ ID n° 86.
 - 4. Utilisation selon l'une quelconque des revendications 1 à 2, dans laquelle le ou lesdits gènes comprennent la

séquence de nucléotides SEQ ID n° 87.

- 5. Utilisation selon l'une quelconque des revendications 1 à 2, dans laquelle le ou lesdits gènes comprennent la séquence de nucléotides SEQ ID n° 94.
- 6. Utilisation selon l'une quelconque des revendications 1 à 2, dans laquelle le ou lesdits gènes comprennent la séquence de nucléotide SEQ ID n° 107.
- 7. Utilisation selon l'une quiconque des revendications 1 à 2, dans laquelle le ou lesdits gènes comprennent les séquences de nucléotides SEQ ID n° 86 et SEQ ID n° 87.
 - 8. Utilisation selon l'une quelconque des revendications 1 à 2, dans laquelle le ou lesdits gènes comprennent les séquences de nucléotides SEQ ID n° 86, SEQ ID n° 87, SEQ ID n° 94 et SEQ ID n° 107.
- 9. Utilisation selon l'une quelconque des revendications 1 à 8, dans laquelle ledit ARN est isolé à partir d'un échantillon avant la détection dudit niveau d'ARN exprimé par ledit gène.
 - 10. Utilisation selon l'une quelconque des revendication 1 à 9, dans laquelle ledit niveau d'ARN est détecté par amplification en chaîne par polymérisa (PCR).
 - 11. Utilisation selon l'une quelconque des revendications 1 à 9, dans laquelle ledit niveau d'ARN est détecté par hybridation.
- 12. Utilisation selon la revendication 11, dans laquelle ledit niveau d'ARN est détecté par hybridation avec und oligonucléotide.
 - 13. Utilisation selon la revendication 12, dans laquelle ledit oligonucléotide comprend de l'ADN, de l'ARN, de l'ADNc, de l'ANP, de l'ADN génomique ou des oligonucléotides de synthèse.

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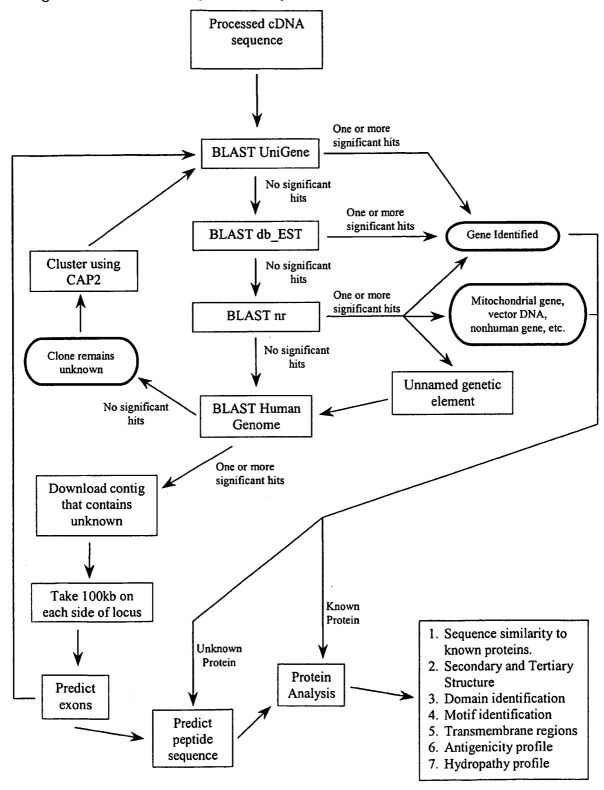


Figure 1: Novel Gene Sequence Analysis

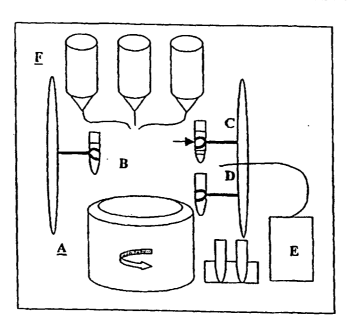


Figure 2. Automated Mononuclear Cell RNA Isolation Device

FIGURE 3

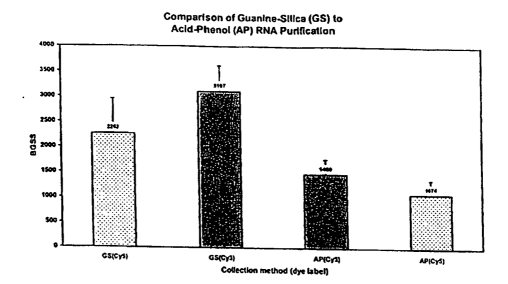


FIGURE 4

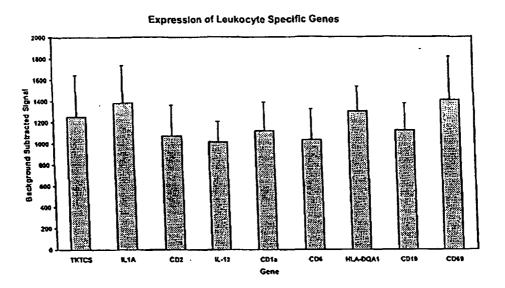
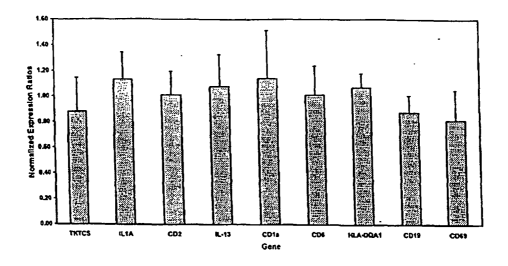


FIGURE 5

Expression of Leukocyte-Specific Genes



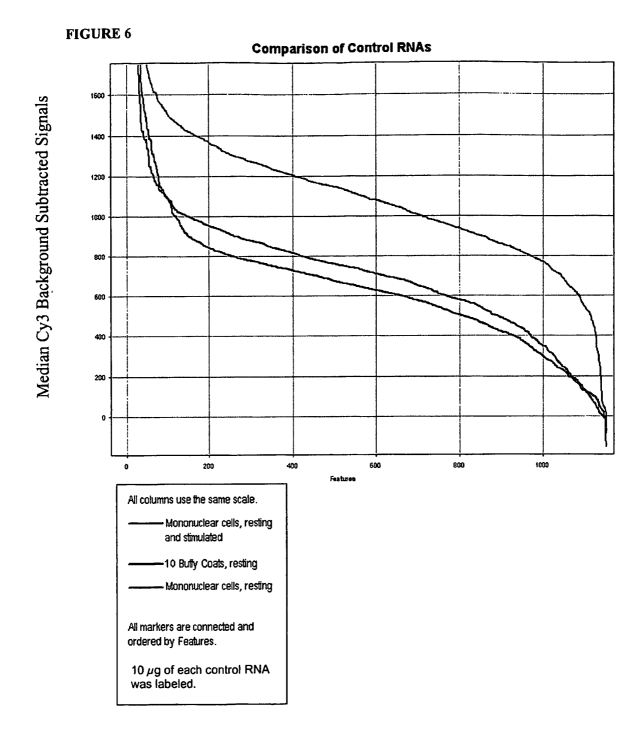
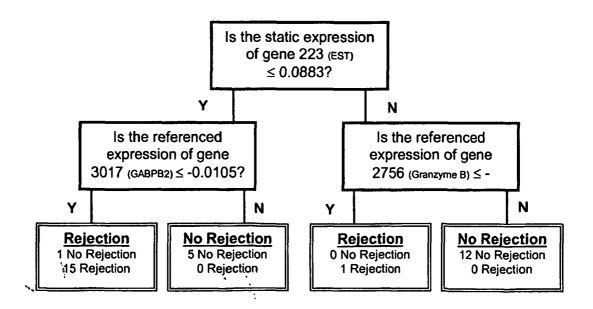


Figure 7: Cardiac Allograft rejection diagnostic genes.

A.

	Marker Gene Expression Ratios					
Sample	Grade	3020	3019	2760	3018	85
12-0025-02	0	3.90	3.69	5.49	3.24	3.34
12-0024-04	0	3.66	4.05	5.89	3.75	3.03
15-0024-01	0	3.55	4.01	5.61	2.90	3.23
12-0029-03	0	3.44	3.12	4.25	3.55	3.07
12-0024-03	0	2.88	2.54	2.56	2.20	2.38
14-0021-05	0	1.31	1.03	1.07	0.91	0.99
14-0005-06	3A	0.42	0.27	0.51	0.22	0.26
14-0012-07	3A	0.60	0.62	0.70	0.42	0.61
14-0001-06	3A	0.93	0.71	0.58	0.37	0.44
14-0009-01	3A	0.71	0.63	0.68	0.61	0.66
12-0012-02	3A	0.86	0.85	0.73	0.41	0.72
12-0001-01	3A	1.08	0.97	1.01	0.40	1.06
Avera	ge Grade 0:	3.13	3.07	4.14	2.76	2.67
Average	Grade 3A:	0.77	0.68	0.70	0.40	0.62
Fold	Difference:	4.08	4.55	5.91	6.82	4.28

B. CART classification model.



C. Surrogates for the CART classification model.

Primary Splitter	static 223	<u>ref 3017</u> ref 4	
Surrogate 1	ref 167	ref 102	ref 2761
Surrogate 2	ref 3016	static 36	ref 2762
Surrogate 3	ref 1760	ref 2764	ref 3016
Surrogate 4	ref 85	ref 2759	ref 2757
Surrogate 5	ref 2763	ref 2761	ref 2758

Figure 8A: Validation of differential expression of Granzyme B in CMV patients using Real-time PCR

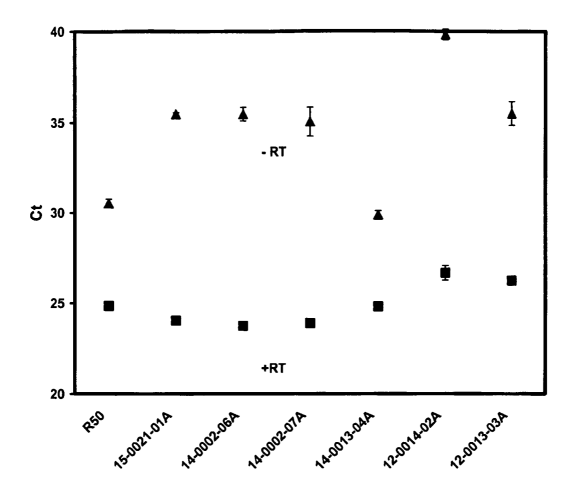


Figure 8B.

QPCR of Granzyme B

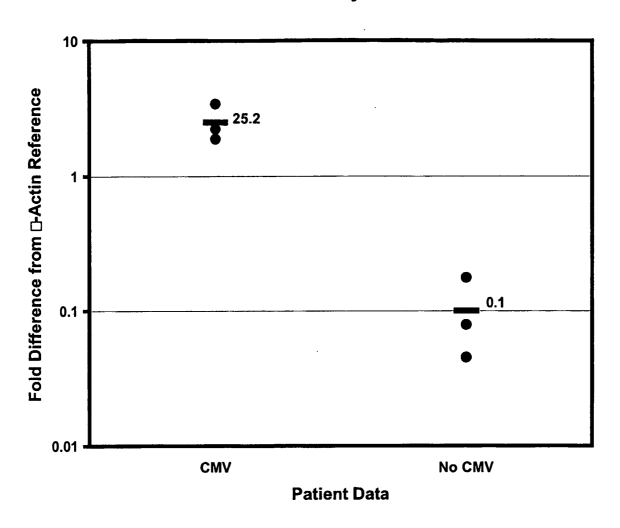


Figure 9

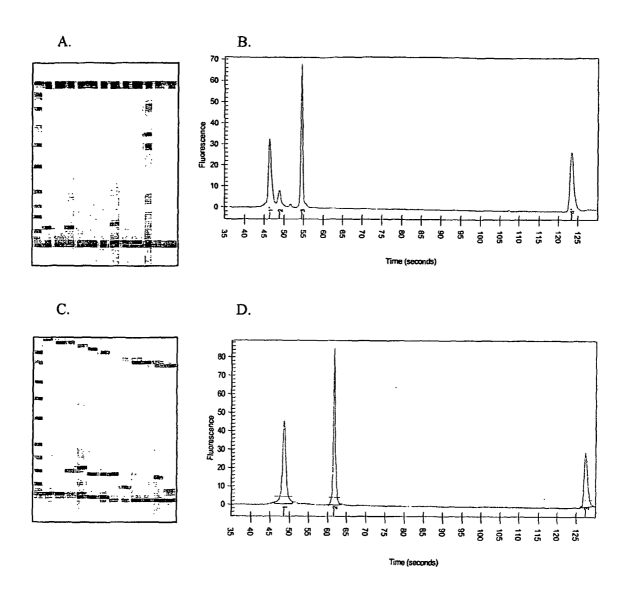


Figure 10

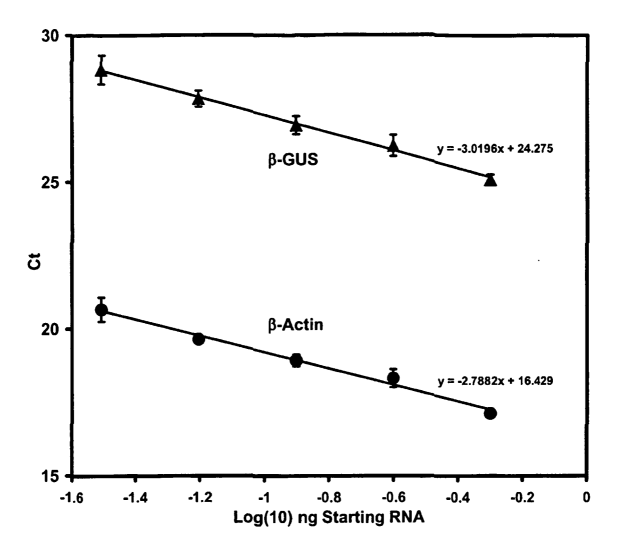
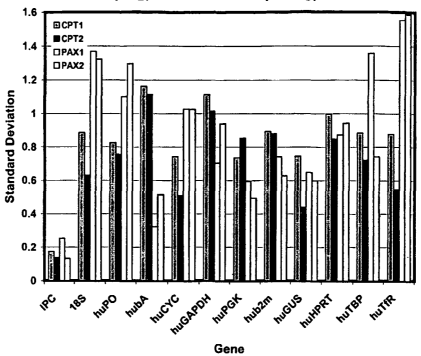


Figure 11





Intensity of Control Genes from PAX RNA (2ug) and CPT RNA (0.5 ug)

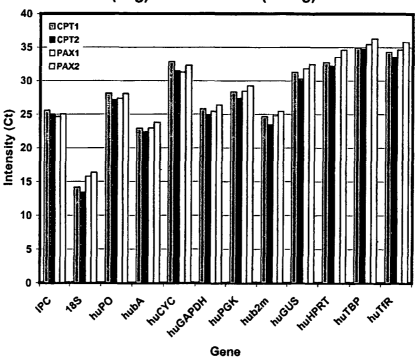


Figure 12

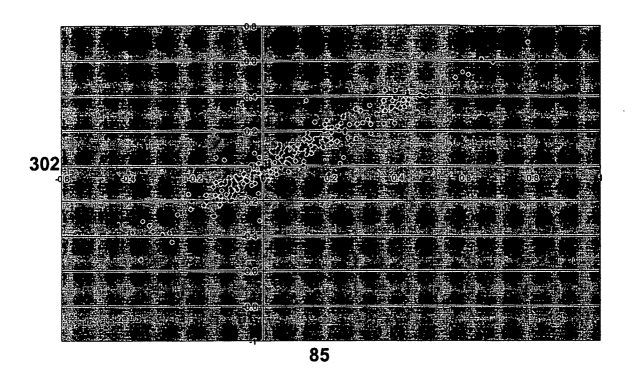
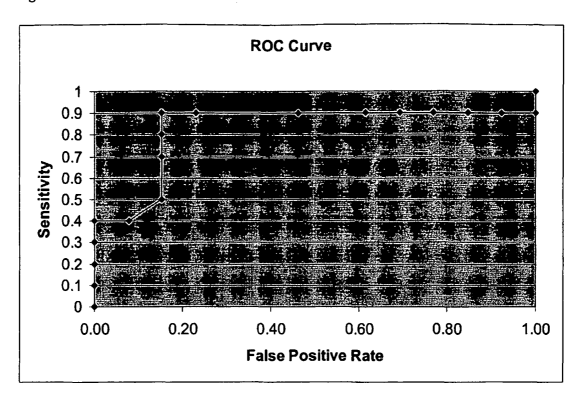


Figure 13



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摘要(译)

描述了通过检测患者中一种或多种基因的表达水平来诊断或监测患者中的移植排斥,特别是心脏移植排斥的方法。还描述了用于诊断或监测移植排斥,特别是心脏移植排斥的诊断寡核苷酸和包含其的试剂盒或系统。

Gene	Cell type
CD10	B-lymphoblasts
RAG1	B-lymphoblasts
RAG2	B-lymphoblasts
NF-E2	Platelets/Megakaryocyte/Erythroid
GATA-1	Platelets/Megakaryocyte
GP IIb	Platelets
pf4	Platelets
EPO-R	Erythroblast
Band 4.1	Erythrocyte
ALAS2	Erythroid specific heme biosynthesis
hemoglobin chains	Erythocyte
2,3-BPG mutase	Erythrocyte
CD16b	Neutrophil
LAP	Neutrophil
CD16	NK cells
CD159a	NK cells