Abstract:

Disclosed is a method for diagnosing melanoma in a human subject, as well as a method for providing a prognosis to a human subject who is at risk of developing melanoma recurrence, and a method for determining the stage of melanoma in a human subject, comprising the step of determining the level of expression of phosphatase and actin regulator 1 (PHACTR1) gene, or fragments thereof, either alone or in combination with the level of expression of secreted integrin-binding phosphoprotein (SPPl), preferentially expressed antigen in melanoma (PRAME), growth differentiation factor 15 (GDF15), and chemokine C-X-C motif ligand 10 (CXCL10) genes. Further, the invention relates to a diagnostic kit, comprising at least one substance for detection of the expression of PHACTR1 or fragments thereof, either alone or in combination with the detection of SPPl, PRAME, GDF15, and CXCL10, for the diagnosis or prognosis of melanoma.
BIOMARKERS FOR DIFFERENTIATING MELANOMA FROM BENIGN NEVUS IN THE SKIN

CROSS-REFERENCE TO RELATED APPLICATIONS

This application claims priority to and benefit of United States Provisional Application No. 61/384,707, filed September 20, 2010, entitled "BIOMARKER FOR DIFFERENTIATING MELANOMA FROM BENIGN NEVUS IN THE SKIN" and United States Provisional Application No. 61/470,682, filed April 1, 2011, entitled "METHODS FOR DIFFERENTIATING MELANOMA FROM BENIGN NEVUS IN THE SKIN".

Each of the aforementioned applications is incorporated herein by reference in its entirety for all purposes.

FIELD OF INVENTION

Provided herein is a method for detection of melanoma by monitoring the presence and levels of specific biomarkers in samples obtained from patients suspected of melanoma. Also provided herein are methods and kits for diagnosing melanoma and providing prognosis of melanoma development in patients based on the presence and levels of specific biomarkers. Further provided herein are methods for determining the stage of melanoma based on the presence of malignant melanoma cells in sentinel lymph nodes of patients.

BACKGROUND OF THE INVENTION

The incidence of cutaneous melanoma has been increasing rapidly over the past several decades, and it is currently the 5th and 7th most common cancer in men and women, respectively, in the US (Jemal, A., Siegel, R., Xu, J. & Ward, E. (2010) Cancer Statistics, CA Cancer J Clin. 60(5):277-300). Although melanoma represents only 4% of all skin cancers, it accounts for 80% of deaths from all skin cancers. Due to the marked difference in five-year survival probability between metastatic (<15%) and localized disease (98%) (Miller, A.J. & Mihm, M.C. Melanoma. (2006) The New England Journal of Medicine 355, 51-65), early detection and diagnosis is the best approach to improve melanoma outcome.

The reason for this diagnostic difficulty is that melanocytes also give rise to a variety of benign neoplasms called melanocytic nevi, which can mimic melanoma histologically. Conversely, some malignant melanomas can mimic benign nevi. Furthermore, the recognition of the many histological features used in diagnosis can be subjective and dependent on experience and thus cause poor inter-observer reproducibility. None of the IHC markers being used today in melanoma diagnosis can differentiate between melanoma and benign nevus (Ugurel, S., Utikal, J. & Becker, J.C. (2009) "Tumor biomarkers in melanoma" Cancer Control: Journal of the


By comparing chromosomal DNA copy number changes between melanomas and benign nevi, four genomic loci (6p25, centromere 6, 6q23, and 11ql3) were selected to develop a multicolor fluorescence in situ hybridization (FISH) assay (Gerami, P. et al. (2009) "Fluorescence in situ hybridization (FISH) as an ancillary diagnostic tool in the diagnosis of melanoma" The American Journal of Surgical Pathology 33, 1146-56). An algorithm based on signal counts (i.e., copy number gains or losses) and percentage of cells with aberrations was shown to have 86.7% sensitivity and 95.4% specificity for melanoma in a validation cohort of unequivocal cases of melanomas and nevi (Gerami, P. et al. (2009) "Fluorescence in situ
hybridization (FISH) as an ancillary diagnostic tool in the diagnosis of melanoma" The American Journal of Surgical Pathology 33, 1146-56). However, in classifying histologically ambiguous lesions, when compared to clinical outcome, the sensitivity and specificity of the FISH probes in identifying malignant diseases were 43% and 80%, respectively, in one study (Vergier, B. et al. (2011) "Fluorescence in situ hybridization, a diagnostic aid in ambiguous melanocytic tumors: European study of 113 cases" Mod. Pathol. 24(5):613-23), and 60% and 50% in another (Gaiser, T. et al. (2010) "Classifying ambiguous melanocytic lesions with FISH and correlation with clinical long-term follow up". Modern Pathology 23, 413-9), raising concerns about their clinical utility in classifying difficult cases. This assay is the only commercially available molecular diagnostic test on the market today (MelanoSite™, Neogenomics/Abbot Molecular).

based 37-gene classifier, which demonstrated 89% concordance with histopathology in differentiating melanoma from benign nevi (Koh, S.S. et al. (2009) "Molecular classification of melanomas and nevi using gene expression microarray signatures and formalin-fixed and paraffin-embedded tissue" Modern Pathology 22, 538-46). Kashani-Sabet and coworkers used the results of their microarray studies to develop a 5-gene (ARPC2, FN1, RGS1, SPP1, and WNT2) immunohistochemistry (IHC) panel, which demonstrated 91% sensitivity and 95% specificity for melanoma (Kashani-Sabet, M. et al. (2009) "A multi-marker assay to distinguish malignant melanomas from benign nevi" Proc. Natl. Acad. Sci. U S A 106, 6268-6272); however, it is important to point out that these values were based on the entire training dataset and thus likely to be significantly overestimated.

These different approaches of translating differential gene expression into clinical diagnostics all have significant limitations. Both RT-PCR and microarray methods require RNA extraction, and the resulting gene expression measurements represent an "average" of heterogeneous groups of cells. Careful microdissection can help, but it is too cumbersome for routine practice. IHC is an attractive approach to provide cellular resolution, but translating differential gene expression at the RNA level to IHC can be difficult, because the best candidate markers may not have suitable antibodies for IHC. This means either a major undertaking to develop new antibodies or use of a suboptimal gene panel based on antibody availability.

Because of the great cellular heterogeneity in a typical melanocyte lesion, RNA in situ hybridization (ISH) is the preferred method for gene expression analysis, since it can be used to determine gene expression at single cell resolution Suzuki, I. & Motokawa, T. (2004) "In situ hybridization: an informative technique for pigment cell researchers” Pigment Cell Research 17, 10-4). However, existing RNA ISH methods have poor sensitivity and specificity and are technically demanding, making them impractical for most of the genes identified in microarray analysis.

A novel RNA ISH technology platform called RNAscope® was recently developed. RNAscope® has single-molecule sensitivity, uses a 6-hour IHC-like procedure, and is multiplex-capable, and compatible with formalin-fixed paraffin-embedded (FFPE) tissue. This makes it possible for the first time to rapidly develop RNA ISH assays for any candidate genes identified from microarrays.
The importance of tumor microenvironment in tumor invasion and metastasis has been firmly established (Hanahan, D. & Weinberg, R.A. (2011) "Hallmarks of Cancer: The Next Generation" Cell 144, 646-674). However, to date cancer diagnostics has either focused exclusively on tumor cell-intrinsic markers (e.g., see Gerami, P. et al. (2009) "Fluorescence in situ hybridization (FISH) as an ancillary diagnostic tool in the diagnosis of melanoma" The American Journal of Surgical Pathology 33, 1146-56) or lack the ability to interrogate tumor and stroma separately in a practical manner (e.g., see Paik, S. et al. (2004) "A multigene assay to predict recurrence of tamoxifen-treated, node-negative breast cancer" The New England Journal of Medicine 351, 2817-26; van't Veer, L.J. et al. (2002) "Gene expression profiling predicts clinical outcome of breast cancer" Nature 415, 530-6; Finak, G. et al. (2008) "Stromal gene expression predicts clinical outcome in breast cancer" Nature Medicine 14, 518-27). For example, a stroma-derived gene expression signature for breast cancer prognosis has been identified before (Finak, G. et al. (2008) "Stromal gene expression predicts clinical outcome in breast cancer" Nature Medicine 14, 518-27). Its implementation as a microarray- or RT-PCR-based test for clinical practice will be challenging since it will require careful microdissection of the stroma from the tumor and it cannot differentiate the many stromal cell types. The role of tumor-host interactions in melanoma progression is well recognized (Hsu, M.-Y., Meier, F. & Herlyn, M. (2002) "Melanoma development and progression: a conspiracy between tumor and host" Differentiation 70, 522-36). Alterations in expression of growth factors, receptors and ECM proteins take place early and work in concert with the accumulative genetic changes in melanocytes to promote melanoma progression. Therefore, the tumor-stroma interface offers abundance of targets for diagnostics, which can be exploited by RNAscope®.

The first RNA ISH-based multi-gene assay for the diagnosis of malignant melanoma and benign nevus was developed in the present invention. This assay determines the expression of PHACTR1 gene either alone or in combination with one or more of SPP1, PRAME, GDF15, and/or CXCL10 genes at cellular resolution. In contrast to earlier approaches noted above, RNAscope® enables a novel biomarker strategy which detects gene expression changes in both the tumor cells and tumor-associated stroma for increased clinical sensitivity and specificity for early stage melanoma. The assays developed in the present invention allow pathologists to seamlessly integrate novel molecular markers with conventional histopathology, significantly advancing today's clinical practice in melanoma diagnosis. Ultimately, the tests result in more
accurate diagnosis of early melanoma while minimizing false positives to spare patients with benign lesions unnecessary treatment and follow-ups.

**SUMMARY OF THE INVENTION**

Therefore, an object of the present invention is to provide new markers for diagnosing the presence of melanoma or for providing a prognosis of the likelihood of recurrence of melanoma, in particular in humans, and for determining the staging of melanoma using such markers.

This object is achieved by providing a method for diagnosing melanoma in a subject, comprising steps of providing a tissue sample from a human subject, wherein the tissue sample comprises a plurality of melanocytes, and determining the level of expression of phosphatase and actin regulator 1 (PHACTR1) marker, or fragments thereof, thereby diagnosing the presence of melanoma based on the expression levels in the provided tissue. In this case, PHACTR1 may be used either alone or, according to another aspect of the invention, in combination with SPP1, PRAME, GDF15, and/or CXCL10 as markers. In yet another aspect of the invention, PHACTR1 may be used in combination with SPP1 as makers.

The object of the invention is fully achieved in this manner. By means of comparative analyses between malignant tissue and non-malignant tissue, genes and proteins can be identified that differ significantly in the types of tissue in their frequency and/or concentration. The quantitative expression of a particular gene or protein in comparison with controls, particularly of the genes and proteins claimed in this invention, thus constitutes an important indicator for the presence of tumor tissues, particularly melanoma, which makes it possible to use these genes and/or proteins as diagnostic markers.

In experiments the inventions is based upon, the inventor discovered that PHACTR1, SPP1, PRAME, GDF15, and/or CXCL10 genes/proteins, in particular PHACTR1 and SPP1 genes/proteins, are well-suited for use as markers for the diagnosis/prognosis of whether or not the tissue sample tested has melanoma.

According to another aspect, the invention concerns a method for diagnosing the presence of melanoma, in which upregulation of PHACTR1, either alone or in combination with upregulation of SPP1, PRAME, GDF15, and/or CXCL10, is indicative of melanoma. In one aspect of the invention, PHACTR1 may be used in combination with SPP1 as makers.
According to one aspect of the invention, determination of the levels of expression of the genes comprises detecting either of the expression level of mRNA expressed form said genes or of the expression level of polypeptides encoded by said genes. In a preferred embodiment, the detection of level of expression of mRNA is performed \textit{in situ}.

In particular, upregulation of the PHACTR1 gene expression is determined, either alone or in combination with the upregulation of SPP1, PRAME, GDF15, and/or CXCL10, in comparison to gene/protein expression of the above genes in normal (or non-malignant) tissue.

In one aspect of the invention, diagnosis of the presence of melanoma is based on the pattern of expression levels of the markers claimed in the present invention. In one embodiment, the level of expression of each marker in the set of markers are determined individually. Based on the pattern of different expression levels of these markers, a diagnosis of the presence of melanoma or a prognosis of the likelihood of the recurrence of melanoma is made. In another embodiment, each marker may be given a weight, based on its power of influence in the analysis. The weighted expression level of the markers are then summed up to produce a score. The score is then used for providing diagnosis or prognosis of melanoma based on a threshold.

In yet another embodiment, the level of expression of each marker in the set of markers are determined indistinctively, \textit{e.g.}, by using an identical label for different markers. The expression level of the markers are pooled and determined altogether. Diagnosis or prognosis of melanoma is then made based on the accumulative measurement.

In addition to diagnosing melanoma and providing prognosis of melanoma by quantitatively measuring the level of expression of the markers claimed in the present invention, a method of qualitatively diagnosing melanoma and providing prognosis of melanoma based on the present or the absence of the markers is also provided in the present invention. In one embodiment, the presence of PHACTR1 marker, either alone or in combination with the presence of SPP1, PRAME, GDF15, and/or CXCL10 markers, is indicative of the presence or the recurrence of melanoma.

The protein sequence of the marker PHACTR1 as well as the gene sequence coding for SPP1, PRAME, GDF15, and CXCL10 are listed in publicly available databases. For example, the mRNA sequences of these genes are included in the database ("GenBank") of the National
Center for Biotechnology Information (NCBI: http://(www).ncbi.nlm.nih.gov/) under database No. NM_030948.1 for PHACTRI, NM_000582.2 for SPPl, NM_006115.3 for PRAME, NM_004864.2 for GDF15, NM_001565 for CXCL10, NM_005511.1 for MLANA, NM_014624.3 for S100A6, and NM_006272.2 for S100B. The gene and protein sequences corresponding to the markers can also be identified via these mRNA numbers, so the present invention expressly include the respective sequences, or fragments thereof, published in the databases and used in the claimed methods.

The mRNA sequences of the diagnostic markers disclosed in the present invention are identified as SEQ ID NO. 1 for PHACTRI, SEQ ID NO. 3 for GDF15, SEQ ID NO. 5 for MLANA, SEQ ID NO. 7 for CXCL10, SEQ ID NO. 9 for PRAME, SEQ ID NO. 11 for S100A6, SEQ ID NO. 13 for S100B, and SEQ ID NO. 15 for SPPl. The protein sequence of the same markers are identified as SEQ ID NO. 2 for PHACTRI, SEQ ID NO. 4 for GDF15, SEQ ID NO. 6 for MLANA, SEQ ID NO. 8 for CXCL10, SEQ ID NO. 10 for PRAME, SEQ ID NO. 12 for S100A6, SEQ ID NO. 14 for S100B, and SEQ ID NO. 16 for SPPl.

The markers can be qualitatively or quantitatively determined at the mRNA level via any gene expression analysis technologies which measure mRNA in solution. Examples of such gene expression analysis technologies include, but not limited to, RNAscope®, RT-PCR, Nanostring, QuantiGene® 2.0, qNPA™, microarray, and sequencing.

In addition to diagnosing the presence of melanoma, the invention also provides a method for differentiating melanoma from benign nevus. According to one aspect of the invention, the presence of PHACTRI markers and one or more of markers selected from the group consisting of SPPl, PRAME, GDF15, and CXCL10 markers is indicative of the presence of melanoma but the absence of benign nevus in the tested human subject. In another aspect of the invention, the presence of PHACTRI and SPPl markers is indicative of the presence of melanoma. In yet another aspect of the invention, the absence of PHACTRI and SPPl markers but the presence of MLANA marker is indicative of benign nevus in the tested human subject.

The invention also concerns a method of determining the stage of melanoma in a human subject. The method comprises the steps of obtaining a tissue sample from a human subject, wherein the tissue sample comprises sentinel lymph node; determining the level of expression of PHACTRI marker, either alone or in combination one or more of SPPl, PRAME, GDF15, and
CXCLIO markers, and determining the stage of melanoma as characterized by the presence of malignant melanoma cells in sentinel lymph node based on the expression levels in the tissue sample. In another embodiment, the method uses a combination of PHACTR1 and SPPl markers.

As mentioned above, in the method of determining the stage of melanoma, determination may be carried out using the level of expression of mRNA expressed from the genes or the level of expression polypeptides encoded by the genes. In addition, a pattern of the level of expression of the markers claimed in the present invention can be used for determining the stage of melanoma. When a pattern of the expression levels is used, the levels of expression of the markers are determined individually.

In the method of determining the stage of melanoma, upregulation of expression levels of markers PHACTR1, SPPl, PRAME, GDF15, and/or CXCLIO predicts recurrence of melanoma. The markers can be qualitatively or quantitatively determined at the mRNA level via any gene expression analysis technologies which measure mRNA in solution, including, but not limited to RNAscope®, RT-PCR, Nanostring, QuantiGene® 2.0, qNPA™, and microarray.

The invention also concerns a diagnostic kit comprising at least one substance for detecting the expression of PHACTR1 marker alone or in combination with one or more of SPPl, PRAME, GDF15, and CXCLIO markers for the diagnosis and prognosis of melanoma. In one embodiment, the diagnostic kit comprises substances for detecting the expression of PHACTR1 and SPPl markers.

This diagnostic kit can be advantageously used to determine the quantity of the markers according to the invention in biological samples, namely by comparing the gene/protein expression of the sample of interest that contains the marker or markers with controls or standards, specifically with respect to increased expression of the marker(s). The diagnostic kit contains e.g. one or more antibodies or oligonucleotides that react with the marker/marker combinations according to the invention at the protein or gene level.

**BRIEF DESCRIPTION OF THE DRAWINGS**

**Fig.1** illustrates the principle of the RNAscope® assay. Step 1: Cells or tissues are fixed and permeablized to allow for target probe access. Step 2: Target RNA-specific oligonucleotide
probes are hybridized to multiple mRNAs. Step 3: Multiple signal amplification molecules that each recognize a specific target probe are hybridized. Each unique label probe is conjugated to a different fluorophore or enzyme. Step 4: Signals are detected using a standard bright-field or epifluorescent microscope.

**Fig. 2** illustrates single copy mRNA detection by RNAscope®. (a) HER2 DNA FISH of HeLa and SK-BR-3 nuclei with RNAscope® target probes and signal amplifiers. Nuclei were co-stained with IL-8 probes as an internal control for diploid gene copy number. Nuclei counterstained with DAPI. (b) Target probes to Her-2 mRNA are hybridized with HeLa cells (left) or SK-BR-3 cells (right). Cells were co-stained with 18S rRNA target probes as an internal control for assay success.

**Fig. 3** illustrates chromogenic detection of mRNA in FFPE by RNAscope®. Breast, lung and prostate FFPE tissue sections hybridized with probes either for UBC (top row, - Figs. 3A, 3C, 3E) or a bacterial gene DapB (bottom row, - Figs. 3B, 3D, 3F). UBC staining was detected in all three tissues; no staining was seen with DapB. Nuclei were counterstained with hematoxylin.

**Fig. 4** illustrates up-regulation of PHACTR1 in melanoma in two microarray datasets. P values were from Kruskal-Wallis test.

**Fig. 5** illustrates detection of mRNAs in melanoma. PHACTR1 mRNA detected as individual red dots. Hematoxylin staining of nuclei (blue) and natural melanin background (brown) are also shown.

**Fig. 6** illustrates an example of staining patterns in a nevus (left, - Figs. 6A, 6B, 6C) and a melanoma (right, - Figs. 6D, 6E, 6F). Positive mRNA staining were detected as red dots. Blue, hematoxylin staining of nuclei.

**Fig. 7** illustrates an example of SPP1 mRNA staining in stromal cells. SPP1. Positive mRNA staining was detected as red dots in scattered stromal cells within the tumor. Blue, hematoxylin staining of nuclei.

**Fig. 8** illustrates RNAscope® staining of a desmoplastic melanoma. The probes for S100A6 and S100B were pooled to detect both together. RNA staining are shown in red. Nuclei are shown in blue.
Fig. 9 illustrates examples of positive staining patterns of GDF15 (Fig. 9A), PRAME (Fig. 9B) and CXCL10 (Fig. 9C) in melanomas. RNA staining is shown in red. Nuclei are shown in blue.

DETAILED DESCRIPTION

Provided herein are based, in part, on the discovery that the presence and level of certain molecules or mRNA in patient samples can be utilized as biomarkers to indicate the presence of myeloma. In particular, these biomarkers can be used to differentiate melanoma from benign nevus in the skin.

1. Definitions

Unless defined otherwise, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art to which the invention pertains. The following definitions supplement those in the art and are directed to the current application and are not to be imputed to any related or unrelated case, e.g., to any commonly owned patent or application. Although any methods and materials similar or equivalent to those described herein can be used in the practice for testing of the present invention, the preferred materials and methods are described herein. Accordingly, the terminology used herein is for the purpose of describing particular embodiments only, and is not intended to be limiting.

The term "polynucleotide" (and the equivalent term "nucleic acid") encompasses any physical string of monomer units that can be corresponded to a string of nucleotides, including a polymer of nucleotides (e.g., a typical DNA or RNA polymer), peptide nucleic acids (PNAs), modified oligonucleotides (e.g., oligonucleotides comprising nucleotides that are not typical to biological RNA or DNA, such as 2'-0-methylated oligonucleotides), and the like. The nucleotides of the polynucleotide can be deoxyribonucleotides, ribonucleotides or nucleotide analogs, can be natural or non-natural, and can be unsubstituted, unmodified, substituted or modified. The nucleotides can be linked by phosphodiester bonds, or by phosphorothioate linkages, methylphosphonate linkages, boranophosphate linkages, or the like. The polynucleotide can additionally comprise non-nucleotide elements such as labels, quenchers, blocking groups, or the like. The polynucleotide can be, e.g., single-stranded or double-stranded.
A "nucleic acid target" or "target nucleic acid" refers to a nucleic acid, or optionally a region thereof, that is to be detected.

A "polynucleotide sequence" or "nucleotide sequence" is a polymer of nucleotides (an oligonucleotide, a DNA, a nucleic acid, etc.) or a character string representing a nucleotide polymer, depending on context. From any specified polynucleotide sequence, either the given nucleic acid or the complementary polynucleotide sequence (e.g., the complementary nucleic acid) can be determined.

The term "gene" is used broadly to refer to any nucleic acid associated with a biological function. Genes typically include coding sequences and/or the regulatory sequences required for expression of such coding sequences. The term gene can apply to a specific genomic sequence, as well as to a cDNA or an mRNA encoded by that genomic sequence.

The term "biomarker" and "marker" as used interchangeably herein, refer to both the protein/gene product in question and the gene coding for this product. The term means a protein, gene product, and/or gene of PHACTR1, SPP1, PRAME, GDF15, and CXCL10.

As used herein the terms "polypeptide" and "protein" as used interchangeably herein, refer to a polymer of amino acids of three or more amino acids in a serial array, linked through peptide bonds. The term "polypeptide" includes proteins, protein fragments, protein analogues, oligopeptides and the like. The term polypeptide as used herein can also refer to a peptide. The amino acids making up the polypeptide may be naturally derived, or may be synthetic. The polypeptide can be purified from a biological sample.

The term "antibody" is used herein in the broadest sense and covers fully assembled antibodies, antibody fragments which retain the ability to specifically bind to the antigen (e.g., Fab, F(ab')2, Fv, and other fragments), single chain antibodies, diabodies, antibody chimeras, hybrid antibodies, bispecific antibodies, humanized antibodies, and the like. The term "antibody" covers both polyclonal and monoclonal antibodies.

The term "biological sample" or "tissue sample" as used herein refers to a sample obtained from a biological subject, including sample of biological tissue or fluid origin, obtained, reached, or collected in vivo or in situ. A biological sample also includes samples from a region of a biological subject containing precancerous or cancer cells or tissues. Such samples can be,
but are not limited to, organs, tissues, fractions and cells isolated from a mammal. Exemplary biological samples include but are not limited to cell lysate, a cell culture, a cell line, a tissue, an organ, an organelle, a biological fluid, and the like. Preferred biological samples include but are not limited to a skin sample, tissue biopsies, and the like.

The term "upregulation" or "increased expression" refers to any gene or protein expression that is increased or enhanced compared to normal expression of the gene in question in healthy samples. Comparison of the differing levels of expression may be carried out using controls or standards. "Upregulated" or "increased" gene expression means that the gene in question is transcribed—in and if applicable, translated into the corresponding protein—to a greater extent than is normal for this gene.

A "label" is a moiety that facilitates detection of a molecule. Common labels in the context of the present invention include fluorescent, luminescent, light-scattering, and/or colorimetric labels. Suitable labels include enzymes and fluorescent moieties, as well as radionuclides, substrates, cofactors, inhibitors, chemiluminescent moieties, magnetic particles, and the like. Patents teaching the use of such labels include U.S. Pat. Nos. 3,817,837; 3,850,752; 3,939,350; 3,996,345; 4,277,437; 4,275,149; and 4,366,241. Many labels are commercially available and can be used in the context of the invention.

A "target probe" is a polynucleotide that is capable of hybridizing to a target nucleic acid and capturing a label probe to that target nucleic acid. The target probe can hybridize directly to the label probe, or it can hybridize to one or more nucleic acids that in turn hybridize to the label probe; for example, the target probe can hybridize to an amplifier or a preamplifier. The target probe thus includes a first polynucleotide sequence that is complementary to a polynucleotide sequence of the target nucleic acid and a second polynucleotide sequence that is complementary to a polynucleotide sequence of the label probe, amplifier, preamplifier, or the like. The target probe is preferably single-stranded.

An "amplifier" is a molecule, typically a polynucleotide, that is capable of hybridizing to multiple label probes. Typically, the amplifier hybridizes to multiple identical label probes. The amplifier also hybridizes to at least one target probe or nucleic acid bound to a target probe. For example, the amplifier can hybridize to at least one target probe and to a plurality of label probes, or to a preamplifier and a plurality of label probes. The amplifier can be, e.g., a linear,
forked, comb-like, or branched nucleic acid. As noted for all polynucleotides, the amplifier can include modified nucleotides and/or nonstandard internucleotide linkages as well as standard deoxyribonucleotides, ribonucleotides, and/or phosphodiester bonds. Suitable amplifiers are described, for example, in USPN 5,635,352, USPN 5,124,246, USPN 5,710,264, and USPN 5,849,481.

A "preamplifier" is a molecule, typically a polynucleotide, that serves as an intermediate between one or more target probes and amplifiers. Typically, the preamplifier hybridizes simultaneously to one or more target probes and to a plurality of amplifiers. Exemplary preamplifiers are described, for example, in USPN 5,635,352 and USPN 5,681,697.

2. Biomarkers

Provided herein are methods relating to the use of mRNA or protein, and other cell marker molecules as biomarkers to detect the presence of melanoma. The presence of biomarkers or their levels can be used to determine whether a given sample has melanoma.

A biological marker or "biomarker" is a substance whose detection indicates a particular biological state, such as, for example, the presence of melanoma. In some embodiments, biomarkers can either be determined individually, or several biomarkers can be measured simultaneously.

In the research that led to the present invention, a plurality of genes have been identified as associating with melanoma. It has furthermore been demonstrated that specific combinations of these genes are associated with melanoma. The plurality of genes associating with melanoma include, but not limited to: PHACTR1 gene (phosphatase and actin regulator 1), SSP1 gene (secreted integrin-binding phosphoprotein), PRAME gene (preferentially expressed antigen in melanoma), GDF15 gene (growth differentiation factor 15), and CXCLIO gene (chemokine C-X-C motif ligand 10). In addition, a plurality of genes have been identified for using as control genes, including: MLANA (melanoma antigen recognized by T-cells), S100A6, S100B, and DapB.
2.1 Selection of PHACTRI, SPP1, and MLANA as Biomarkers for Detecting Melanoma

In one example of the invention, it was found by the inventors of the present invention that PHACTRI and SPP1 are the strongest markers for melanoma. PHACTRI and SPP1 genes were detected at the early phase of melanoma.

In the research that led to the present invention, the gene expression profile of a large number of genes from skin samples ranging from normal skin to benign nevus to melanoma was analyzed. In the samples with abnormal skin condition, the lesions on the skin represented different stages of melanoma progression according to a current model. The model posits that melanoma initiates from dysplastic nevi, which then progresses to melanoma in situ or minimally invasive melanoma (RGP) to invasive melanoma (VGP) to metastatic melanoma (MTS) in a stepwise fashion (Hsu, M.-Y., Meier, F. & Herlyn, M. (2002) "Melanoma development and progression: a conspiracy between tumor and host." Differentiation 70(9-10):522-36). The microarray data of genome-wide gene expression analysis were retrieved from the Gene Expression Omnibus (GEO) database (http://(www.)ncbi.nlm.nih.gov/geo/). The GEO database stores gene expression microarray datasets contributed by various research groups, often after bioinformatics and statistical analysis have been conducted on these datasets by the original authors. In order to identify consistent, cross-validated candidate markers, datasets from two different microarray platforms, GSE3189 (Affymetrix® GeneChip® HG-U133A, 22,000 genes) and GSE12391 (Agilent® Human Whole Genome Oligo Microarray, 41,000 genes), were chosen by the inventors for candidate identification.

In the present invention, it has been surprisingly found that increased expression level of PHACTRI gene indicates skin samples having melanoma. Before the present invention, PHACTRI gene has never been selected as biomarker for detection or prognosis of melanoma in any studies. In the present invention, PHACTRI gene was identified as biomarker for melanoma based on the result of novel statistical algorithm and classification method. Limma linear models (Smyth, G.K. Limma (2005) "Linear models for microarray data. Bioinformatics and Computational Biology Solutions using R and Bioconductor" R. Gentleman, V. Carey, S. Dudoit, R. Irizarry, W. Huber (eds.), Springer, New York, pages 397^-20) for microarrays was used to find genes differentially expressed between normal skin/benign nevi and melanomas. Limma linear models were developed by Smyth G. K. Limma. This algorithm uses empirical
Bayes shrinkage of gene-wise residuals to ensure stable linear models when the number of samples is small, as is the case in the present invention. Using the Limma linear models, PHACTR1 was identified as the most strongly up-regulated gene in microarray GSE12391 and the second most strongly up-regulated gene in melanoma in microarray GSE3189 (see Table 1). The p values were adjusted after controlling for multiple hypothesis testing. PHACTR1 demonstrated up-regulation in the early stage of minimally invasive melanoma, whereas its expression in dysplastic nevi was similar to that in common nevi (Fig. 4), suggesting its utility to differentiate between dysplastic nevi and melanomas.

Another aspect of the invention concerns the use of expression level of PHACTR1 and SPP1 markers to predict whether a skin sample has melanoma. It was surprisingly found that increased expression level of SPP1 marker in combination with increased expression level of PHACTR1 marker indicate the presence of melanoma in the skin samples tested. Thus, the SPP1 gene is a second marker for diagnosing melanoma. Together, the PHACTR1 gene and SPP1 gene, are the strongest markers for melanoma and were detected at the early phase of melanoma. It was found in the present invention that, as shown in Table 2, among the 78 melanoma samples tested, 83% and 91% of the melanomas were positive for PHACTR1 and SPP1, respectively, whereas none of the 19 nevi tested had detectable signals for either marker. In combination, 95% of the melanomas were positive for at least one marker: 84% were positive for both, 12% and 4% were positive for either SPP1 or PHACTR1 alone, respectively.

The myriad functions of SPP1 include cell-ECM adhesion as a ligand for alpha (-v) -beta containing integrins and CD44 and as an activator of matrix metalloproteases (MMP2 in particular) for ECM remodeling, both of which are important in tumor invasion. The PHACTRI protein is a member of the PHACTR/scapinin family, which binds to both protein phosphatase 1 (PP1) and actin. Expression of PHACTR3 enhances cell spreading and motility through its PP1-binding and actin-binding activities (Sagara, J., Arata, T. & Taniguchi, S. (2009) "Scapinin, the protein phosphatase 1 binding protein, enhances cell spreading and motility by interacting with the actin cytoskeleton" PLoS One 4, e4247).

Without being limited by theory, it is likely that PHACTRI may function to regulate tumor invasion and migration. Comparing to benign nevi, a defining hallmark of malignant melanoma is the ability to invade into the dermis and metastasize to distant sites. Therefore, the mechanistic involvement in tumor invasion and metastasis by SPP1 and PHACTRI may explain their utility as markers for discriminating melanoma from benign nevi. There has not yet been any connection described between the expression level of PHACTRI and the diagnosis/prognosis of melanoma. There was also no connection described between the expression level of a set of markers consisting of PHACTRI and SPP1 and the diagnosis/prognosis of melanoma. There was no teaching of a in situ hybridization assay using a combination of PHACTRI and SPP1 markers for the diagnosis/prognosis of melanoma.

In yet another example of the invention, the present invention also concerns the use of MLANA gene as a positive control for the detection of melanoma. MLANA gene expresses in both normal melanocytes and malignant melanocytes such as melanoma cells but not in other type of cells in the skin. Thus, in one embodiment, MLANA was used as a biomarker for cell type identification. The presence of MLANA is indicative of the presence of either normal melanocytes or melanoma cells or both. The absence of MLANA is indicative the absence of either normal melanocytes or melanoma cells. In one embodiment, if a sample tissue is detected to have the presence of MLANA marker but the absence of PHACTRI and SPP1 markers, the sample tissue is considered to have benign nevus.

MLANA is also used in the present invention as a positive control for assay success. Very often detection of markers is performed on Formalin-Fixed, Paraffin-Embedded (FFPE) tissue microarrays (TMAs). These TMAs were often constructed from archival tissues from
various times and sources, and thus could vary in the quality of RNA preservation. If MLANA can be detected in a FFPE tissue sample, it indicates that the sample must have acceptable quality. Thus, MLANA also serves as a positive control for RNA integrity.

2.2 Detection of Melanoma Using Biomarkers of PHACTRI, SPPl and/or MLANA

One aspect of the invention provides methods for detection of melanoma cells in a sample based on the expression level of PHACTRI, SPPl and MLANA markers. In one embodiment, upregulated level of expression of PHACTRI and SPPl markers is indicative of the presence of melanoma. Detection of changes in the level of expression of markers often depends on the sensitivity of the assay used. In an assay with sufficiently high sensitivity, the presence of PHACTRI and SPPl markers is indicative of the presence of melanoma. Samples tested with unchanged level of or in absence of PHACTRI and SPPl markers but with upregulated level or with the presence of MLANA marker are considered to have benign nevus. Samples tested upregulated or with the presence of either PHACTRI marker or SPPl marker or both markers, and also upregulated or with the presence of MLANA marker are considered to have melanoma. MLANA gene serves as positive control for assay success and a marker for melanocytes. In addition, bacterial gene DapB is used as negative control. For example, the expression of PHACTRI and SPPl may be determined as positive or negative using the DapB staining as a threshold.

In one embodiment, provided herein is a method for diagnosing melanoma in a human subject suspected of melanoma, comprising the following steps:

(a) obtaining a tissue sample from said human subject, wherein said tissue sample comprises a plurality of melanocytes;

(b) determining the level of expression of PHACTRI and SPPl genes, or fragments thereof; and

thereby diagnosing the presence of melanoma based on the expression levels in said tissue sample.

In another embodiment, provided herein is a method for detecting melanoma in a human subject, wherein the presence of either PHACTRI or SPPl gene is indicative of melanoma in said human subject.
In another embodiment, the detecting step of the above method for further comprising
detecting the present of a MLANA gene, wherein the presence of MLANA gene is indicative of
melanoma and benign nevus, and the absence of MLANA gene is indicative of other cell types.

In addition to the initial prediction of the presence or absence of melanoma in a sample of
a patient, melanoma can be further differentiated from benign nevus by monitoring the presence
and the absence of PHACTRI gene and SPPI gene, optionally with MLANA gene serving as
positive control.

In one embodiment, provided herein is a method for differentiating melanoma from
benign nevus in a human subject, wherein the presence of either PHACTRI or SPPI gene is
indicative of melanoma in said human subject, and the absence of both PHACTRI and SPPI
gene is indicative of benign nevus in said human subject.

In one exemplary embodiment illustrated in Fig. 6, skin samples of a human subject with
benign nevus and skin samples of human subject with melanoma were tested for the presence of
marker genes PHACTRI, SPPI and MLANA. In skin sample of presented in Figs. 6A, 6B and
6C, the sample is completely negative for PHACTRI and SPPI, and positive for MLANA,
demonstrating that the sample is absent of melanoma cells but have melanocytes and thus has
benign nevus. In the sample presented in Figs. 6D, 6E and 6F, the sample is positive for all three
genes: PHACTRI, SPPI and MLANA, indicating it has melanoma, which is a malignant type of
melanocytes.

In one embodiment of the present invention, the pattern of expression levels of marker
genes PHACTRI and SPPI are used to diagnose the presence of melanoma or to provide
prognosis of the likelihood of recurrence of melanoma. In one embodiment, the levels of
expression of PHACTRI and SPPI are determined individually, e.g., using distinctive labels for
each gene. The pattern of levels of expression of PHACTRI and SPPI genes are then analyzed
to determine the presence or absence of melanoma in the tested sample. In another embodiment,
levels of expression of PHACTRI and SPPI genes are given different weights based on their
degree of influence in determining the presence of absence of melanoma. In another
embodiment, the levels of expression of PHACTRI and SPPI are pooled and determined
altogether, e.g., using an identical label for all genes. The combined expression level is then
analyzed to determine the presence or absence of melanoma in the tested sample.
The present invention additionally identified two types of SPP1 staining pattern. In one type of SPP1 staining pattern, as shown in Fig. 7, the staining of SPP1 was detected predominantly in tumor-associated macrophages (TAMs), especially at the interface between tumor and stroma. Another type of SPP1 staining pattern is shown in Fig. 6F (SPP1 in melanoma). The staining is in both TAMs and tumor cells, and thus appears scattered around in the sample.

2.3 Selection of PRAME, GDF15, CXCL10, S100A6 and S100B Genes as Additional Biomarkers for Improved Detection of Melanoma

As mentioned in Section 2.1 above, it was found in the present invention that a combination of PHACTR1 and SPP1 genes can detect 95 % of the melanomas. However, PHACTR1 an SPP1 genes are negative for the remaining 5% of the melanomas. Further, samples detected positive for PHACTR1 and SPP1 genes must also be positive for the expression of MLANA gene as the later is used a positive control for the detection.

Desmoplastic melanoma represents 2-3% of all melanoma. However, desmoplastic melanoma is negative in expression of MLANA. Therefore, in order to improve the scope of the melanoma detection, identification of additional marker genes which can be used to improve the scope of the melanoma detection was performed in the present invention.

Soikkeli, J. et al. (2007) "Systematic search for the best gene expression markers for melanoma micrometastasis detection" The Journal of Pathology 213, 180-9; Westekemper, H. et al. (2010) "Expression of MCSP and PRAME in conjunctival melanoma" British Journal of Ophthalmology 94, 1322-1327; Riker, A.I. et al. (2008) "The gene expression profiles of primary and metastatic melanoma yields a transition point of tumor progression and metastasis" BMC Medical Genomics 1, 13; Amatschek, S. et al. (2011) "CXCL9 induces chemotaxis, chemorepulsion and endothelial barrier disruption through CXCR3-mediated activation of melanoma cells" British Journal of Cancer 104, 469-79; Khaitan, D. et al. (2011) "The Melanoma-Upregulated Long Noncoding RNA SPRY4-IT1 Modulates Apoptosis and Invasion" Cancer Research 71(1):3852-62. RNAseq assays for the 9 selected genes were performed to test their diagnostic performance in FFPE tissue microarrays of 100 samples including 77 melanoma and 23 benign nevus samples. Among the 9 candidate genes, three genes, PRAME, GDF15, CXCLIO, were selected for their superb specificity to be expressed only in melanoma but not in benign nevi. These three genes all show positive staining in melanomas and negative staining in nevi, with no false positives (Fig. 9, and Table 3). Before the present invention, there was no connection described between the expression level of a gene set consisting of PHACTR1, SPP1, PRAME, GDF15, and CXCLIO or any combination thereof and the diagnosis/prognosis of melanoma.

S100A6 and S100B were known for expression in desmoplastic melanoma which are normally melan-A-negative (Fig. 8). Thus, S100A6 and S100B genes serve as control for assay success and also as positive control for desmoplastic melanoma.

2.4 Detection of Melanoma Using Biomarkers of PHACTR1, SPP1, MLANA, PRAME, GDF15, CXCLIO, S100A6 and/or S100B

One aspect of the invention provides methods for detection of melanoma cells in a sample based on the level of expression of PHACTR1, SPP1, MLANA, PRAME, GDF15, CXCLIO, S100A6 and S100B markers. In one embodiment, samples tested upregulated for any one of PHACTR1, SPP1, PRAME, GDF15, and CXCLIO markers are considered to have melanoma. In a specific embodiment, samples tested negative (absence) for PHACTR1, SPP1, PRAME, GDF15, and CXCLIO markers but positive (presence) of MLANA, S100A6 or S100B genes are considered to have benign nevus. Samples tested positive for any one of PHACTR1,
SPPl, PRAME, GDF15, and CXCLIO markers or any combination thereof, together with positive result for any one of MLANA, S100A6 and S100B markers or any combination thereof are considered to have melanoma. MLANA, S100A6 and S100B gene serve as positive control for assay success and a marker for melanocytes. In addition, bacterial gene DapB serves as negative control.

In one embodiment, provided herein is a method for detecting melanoma in a human subject having a skin lesion suspected of melanoma, comprising the following steps:

(a) obtaining a tissue sample of said skin lesion from said human subject, wherein said tissue sample comprises a plurality of epidermal cells; and

(b) detecting the presence of a plurality of genes in said tissue sample, wherein said plurality of genes comprise PHACTRI, SPPl, PRAME, GDF15, and CXCLIO markers, wherein the presence of any one of the five gene is indicative of melanoma in said human subject.

In another embodiment, the detecting step of the above method for further comprising detecting the present of MLANA, S100A6 and S100B markers. The presence of MLANA, S100A6, or S100B markers is indicative of melanoma or and benign nevus, and the absence of these genes is indicative of other cell types.

In addition to the initial prediction of the presence of melanoma in a sample of a patient, the present invention also provides a method for differentiating melanoma from benign nevus based on the expression level of PHACTRI, SPPl, PRAME, GDF15, and CXCLIO genes, with MLANA, S100A6 and S100B genes serving as positive controls.

In one embodiment, provided herein is a method for differentiating melanoma from benign nevus in a human subject, comprising the following steps:

(a) obtaining a tissue sample of said human subject, wherein said tissue sample comprises a plurality of melanocytes; and

(b) detecting the presence of a plurality of genes in said tissue sample, wherein said plurality of genes comprise PHACTRI marker, either alone or in combination with any one or more of SPPl, PRAME, GDF15, and CXCLIO markers,
wherein the presence of any one of the five genes is indicative of melanoma in said human subject, and the absence of both all five gene is indicative of benign nevus in said human subject.

In another embodiment, the detecting step of the above method for further comprising detecting the presence of MLANA, S100A6 and S100B genes. The presence of MLANA, S100A6 or S100B genes is indicative of melanoma or and benign nevus, and the absence of these genes is indicative of other cell types.

2.5 Measuring The Expression Levels of PHACTR1, SPP1, PRAME, GDF15, and CXCL10 Genes

The invention also concerns the determination of the expression level of PHACTR1 maker, and optionally in combination with the expression level of SPP1, PRAME, GDF15, and CXCL10 markers in a tissue sample. In a preferred embodiment, the determination is made in situ using a gene expression analysis assay which measures mRNA in solution. According to one aspect, the sample may be a solid body sample from a mammal, especially a sample of skin. The sample may be taken and if necessary prepared according to methods known to a person skilled in the art. As will be disclosed below, in the method according to the invention, detection/determination of gene expression level may be carried out using the auxiliary methods of RNAscope®, Nanostring, QuantiGene® 2.0, qNPA™ ELISA (enzyme-linked immunosorbent assay), gene/protein microarrays, RT-PCR, sequencing, etc. These methods are known in prior art, particularly in the area of medical research. Instructions for the use of some these methods and other methods not discussed here may be found e.g. in Sambrook, Fritsch, and Maniatis, "Molecular Cloning, A Laboratory Manual".

In one embodiment, provided herein is a method for measuring a plurality of markers in a human subject, comprising the following steps:

(a) obtaining a tissue sample from said human subject, wherein said tissue sample comprises a plurality of melanocytes; and

(b) determining the level of expression of PHACTR1, either alone or in combination with any one or more of SPP1, PRAME, GDF15, and CXCL10 genes, or fragments thereof.
3. Detection Methods

All aspects of this invention are generally applicable to in situ detection of nucleic acids in individual cells. However, many features of this invention can also be used in whole-sample nucleic acid detection applications.

In a preferred embodiment of the invention, the level of the biological marker is determined in vitro in a biological sample obtained from an individual. For in vitro determining the level of the biological markers of the present invention, any suitable biological sample from bodily skin that may comprise a biological marker identified by this research may be used. The biological sample may be processed according to well-known techniques to prepare the sample for testing.

In diagnosing or identifying a melanoma, a test sample containing at least one cell from melanoma is provided to obtain a genetic sample. The test sample may be obtained using any technique known in the art including biopsy, surgical excisions, needle biopsy, scraping, etc. From the test sample is obtained a genetic sample. The genetic sample comprises a nucleic acid, preferably RNA and/or DNA. For example, in determining the expression of marker genes one can obtain mRNA from the test sample, and the mRNA may be reverse transcribed into cDNA for further analysis. In another embodiment, the mRNA itself is used in determining the expression of marker genes. In some embodiments, the expressions level of a particular marker gene may be determined by determining the level/presence of a gene product (e.g., protein) thereby eliminating the need to obtain a genetic sample from the test sample.

Once a genetic sample has been obtained, it can be analyzed for the presence or absence of particular marker genes. For detection of biological markers of the invention, gene signal detection can be made of conventional methods known in the art. including, but not limited to, RNAscope®, Nanostring, QuantiGene® 2.0, qNPA™, sequencing, PCR, RT-PCR, quantitative PCR, restriction fragment length polymorphism, hybridization techniques, Northern blot, microarray technology, DNA microarray technology, etc. In determining the expression level of a marker gene or genes in a genetic sample, the level of expression may be normalized by comparison to the expression of another gene such as a well known, well characterized gene or a housekeeping gene.
The expression data from a particular marker gene or group of marker genes may be analyzed using statistical methods known in the art in order to determine the phenotype or characteristic of a particular tumor or cancer. Methods used in classifying tumors include, but not limited to: decision trees, support vector machine (SVM), neural network, etc.

3.1 Nucleic Acid Detection Methods

When the biological marker is a gene, and/or a polynucleotide fragment and/or variant thereof, e.g. DNA, cDNA, RNA, mRNA etc., such as a gene coding for a specific protein, or mRNA that is transcribed, the biological marker can be measured in e.g. skin biopsies, by e.g. well-known molecular biological assays, such as in situ hybridization techniques using probes directed to the specific polynucleotides. In one embodiment, the RNAscope® technology may be used to determine the gene expression in situ in FFPE tissue sections of melanocyte lesions. Other nucleic-acid based assays which may be used according to the invention include RT-PCR, nanostring, sequencing, nucleic acid based ELISA, Northern blotting etc, and any combinations thereof.

Generally, a method for detecting a nucleic acid marker comprises hybridizing an oligonucleotide to a marker within the nucleic acid target in a sample from a subject under moderate to high stringency conditions and detecting hybridization of the oligonucleotide using a detection means, such as for example, an amplification reaction or a hybridization reaction or a combination of both.

In another embodiment, a marker that is associated with melanoma occurs within a protein coding region of a genomic gene (e.g. PHACTR1, SPP1, FRAME, GDF15, CXCL10, etc.) and is detectable in mRNA encoded by that gene. Such a marker may be detected using, for example, RNAscope®, nanostring, reverse-transcriptase PCR (RT-PCR), sequencing, transcription mediated amplification (TMA) or nucleic acid sequence based amplification (NASBA), although any mRNA or cDNA based hybridization and/or amplification protocol is clearly amenable to the instant invention.

3.1.1 Nucleic Acid Detection Method Based on RNAscope®

The inventors of this application have developed an in situ hybridization method (United States Patent Application No. 7,709,198, incorporated herein by reference in its entirety) called
RNAscope®, that allows for the direct visualization of RNA in situ. This method utilizes the oligonucleotide probe sets and novel signal amplification systems described below. The assay can be used on a variety of sample types including cultured cells, peripheral blood mononuclear cells (PBMCs), frozen tissue, and formalin-fixed paraffin embedded (FFPE) tissue. In addition, the assay can utilize both chromogenic and fluorescent detection reagents.

One embodiment of this invention concerns the adaptation of RNAscope® assay technology to detect, in situ, marker genes of melanoma in skin sample of a patient.

3.1.1.1 RNAscope® Assay Technology

The RNAscope® assay technology provides multiplex nucleic acid assays in single cells (see Fig. 1). At the core of this technology is the "double Z" probe design, which allows robust amplification of specific hybridization signals without also amplifying nonspecific events. Each target probe ("Z") has a target-specific sequence, which binds to the target mRNA, a spacer, and a "tail" sequence. Two target probes (double Z) hybridize contiguously onto a target mRNA, and the two "tail" sequences form a 28-base hybridization site for the PreAmplifier. The double Z probe design ensures high fidelity of signal amplification because 1) it is highly unlikely that a pair of target probes will hybridize nonspecifically juxtaposed to each other to form a binding site for the PreAmplifier; and 2) neither tail alone can bind efficiently to the PreAmplifier under the assay conditions. The PreAmplifier, Amplifier and Label Probe are hybridized sequentially to each target probe pair, resulting in the accumulation of as many as 8,000 label molecules per 1 kb of target RNA. The Label Probe can be conjugated to either a fluorophore or a chromogenic enzyme (e.g., HRP), enabling viewing of hybridization signals under a standard bright-field or epifluorescent microscope, respectively. With a fluorescent Label Probe, the signals can contain at least 100-fold more fluorescent molecules than traditional RNA fluorescent ISH methods and are readily visible under a standard fluorescent microscope.

In addition, multiple signal amplifiers have been built that each recognizes a unique tail sequence on the target probes, allowing for the simultaneous visualization of multiple target RNAs. Importantly, this assay is compatible with partially degraded RNA present in archival FFPE tissues, since the double Z probe pairs target short regions of -50 nucleotides in length.
3.1.1.2 Single Molecule Detection by RNAscope®

The marked improvement in signal-to-noise ratio with RNAscope® allows detection of single RNA molecules as punctuate dots on the stained slides. To demonstrate this, a probe set was used to detect human epidermal growth factor receptor 2 (HER2) mRNA and genomic DNA in SK-BR-3 and HeLa cells. For genomic DNA, two fluorescent dots were detected in HeLa cells whereas many more dots were seen in SK-BR-3 cells, which is consistent with the HER2 gene amplification present in this cell line (Fig. 2a). The same probe set was used to detect HER2 mRNA transcripts, and the expression levels in HeLa and SK-BR-3 cells were consistent with their gene amplification status (Fig. 2b). These results demonstrate that RNAscope® is capable of single RNA molecule detection. Furthermore, the number of HER2 mRNA dots in HeLa cells were counted and compared that with the number of mRNA transcripts determined by QuantiGene® 2.0, a solution-based assay that can measure RNA transcripts quantitatively. The average copy number per cell determined by RNAscope® was comparable to that determined by QuantiGene® 2.0, again indicating that each dot is likely to represent a single mRNA molecule.

3.1.1.3 Detection of mRNA in FFPE tumor tissues by RNAscope®

Since each double Z probe pair targets a region of -50 bases in length, RNAscope® has built-in robustness against partial RNA degradation, as is often the case in archival FFPE tissue. This was demonstrated by hybridizing probe sets targeting the housekeeping gene ubiquitin C (UBC) to archival breast, lung and prostate tissues following standard tissue pretreatment in boiling citrate buffer and protease digestion. Strong positive staining was detected in all three tissue types, but no signal was present when the negative control DapB probes were used (Fig. 3).

3.1.1.4 Using RNAscope® for Detection of Melanoma

In one example, RNAscope® was used to detect three nucleic acid targets PHACTR1, SPP1, and MLANA in a melanocyte of a patient. In the methods, a sample comprising the cell is provided. The cell tested comprises, or is suspected of comprising, PHACTR1, SPP1, and MLANA. Provided in the assay are: a first label probe comprising a first label, a second label probe comprising a second label, and a third label probe comprising a third label, wherein the signals from the three labels are distinguishable from each other. Alternatively, PHACTR1 and
SPP1 labels are identical and the signals emitted from the two markers are indistinguishable from each other. For each of the three nucleic acid targets PHACTRI, SPP1, and MLANA, three target probe sets, each comprising at least two target probes are provided.

In one embodiment of the present invention, the first nucleic acid target is PHACTRI, the second nucleic acid target is SPP1 and the third nucleic acid target is MLANA. When identical labels are used for PHACTRI and SPP1, detection of label signals in a sample indicates that either PHACTRI or SPP1, or both existed in the sample. The detection of uniformed label signals suggests the presence of melanoma in the sample. When the three labels are distinguishable, detection of each individual labels provides further information regarding the expression of the individual genes in the sample.

The first target probe set is hybridized, in the cell, to the first nucleic acid target PHACTRI (when the first nucleic acid target is present in the cell), the second target probe set is hybridized, in the cell, to the second nucleic acid target SPP1 (when the second nucleic acid target is present in the cell), and the third target probe set is hybridized, in the cell, to the third nucleic acid target MLANA (when the third nucleic acid target is present in the cell). The first label probe is captured to the first target probe set, the second label probe is captured to the second target probe set, and the third label probe is captured to the third target probe set, thereby capturing the first label probe to the first nucleic acid target PHACTRI, the second label probe to the second nucleic acid target SPP1 and the third label probe to the third nucleic acid target MLANA. The first signal from the first label, the second signal from the second label and the third signal from the third label are then detected. Since the first, second and third labels are associated with their respective nucleic acid targets through the target probes, presence of the label(s) in the cell indicates the presence of the corresponding nucleic acid target(s) in the cell. The methods are optionally quantitative. Thus, an intensity of the first, second and third signal can be measured, and the intensity of the first signal can be correlated with a quantity of PHACTRI, SPP1 and MLANA in the cell. As another example, a signal spot can be counted for each copy of the PHACTRI, SPP1 and MLANA genes to quantitate them.

In one aspect, the label probes bind directly to the target probes. In another aspect, the label probes are captured to the target probes indirectly, for example, through binding of preamplifiers and/or amplifiers. Use of amplifiers and preamplifiers can be advantageous in
increasing signal strength, since they can facilitate binding of large numbers of label probes to each nucleic acid target.

In the above classes of embodiments, one target probe hybridizes to each label probe, amplifier, or preamplifier. In alternative classes of related embodiments, two or more target probes hybridize to the label probe, amplifier, or preamplifier.

In embodiments in which two or more target probes are employed, the target probes preferably hybridize to nonoverlapping polynucleotide sequences in their respective nucleic acid target. The target probes can, but need not, cover a contiguous region of the nucleic acid target. Blocking probes, polynucleotides which hybridize to regions of the nucleic acid target not occupied by target probes, are optionally provided and hybridized to the target. For a given nucleic acid target, the corresponding target probes and blocking probes are preferably complementary to physically distinct, nonoverlapping sequences in the nucleic acid target, which nonoverlapping sequences are preferably, but not necessarily, contiguous. Having the target probes and optional blocking probes be contiguous with each other can in some embodiments enhance hybridization strength, remove secondary structure, and ensure more consistent and reproducible signal.

As noted, the methods are useful for multiplex detection of nucleic acids, including simultaneous detection of more than three nucleic acid targets. Thus, the cell optionally comprises or is suspected of comprising a fourth, fifth, sixth, seventh or even more nucleic acid target. The method detect a fourth nucleic acid target comprises: providing a fourth label probe comprising a fourth label, wherein a fourth signal from the fourth label can be either distinguishable or indistinguishable from the first three signals depending the degree of information to be collected, providing at least two fourth target probe, hybridizing in the cell the fourth target probe to the fourth nucleic acid target (when the fourth target is present in the cell), capturing the fourth label probe to the fourth target probe, and detecting the fourth signal from the fourth label. In the present invention, the additional nucleic acid targets include but not limited to PRAME, GDF15, CXCL10, S100A6, S100B, etc.

In detection of nucleic acid targets in a cell, the cell is typically fixed and permeabilized before hybridization of the target probes, to retain the nucleic acid targets in the cell and to permit the target probes, label probes, etc. to enter the cell. The cell is optionally washed to
remove materials not captured to one of the nucleic acid targets. The cell can be washed after any of various steps, for example, after hybridization of the target probes to the nucleic acid targets to remove unbound target probes, after hybridization of the preamplifiers, amplifiers, and/or label probes to the target probes, and/or the like.

The various capture and hybridization steps can be performed simultaneously or sequentially, in essentially any convenient order. Preferably, a given hybridization step is accomplished for all of the nucleic acid targets at the same time. For example, all the target probes (first, second, etc.) can be added to the cell at once and permitted to hybridize to their corresponding targets, the cell can be washed, amplifiers (first, second, etc.) can be hybridized to the corresponding target probes, the cell can be washed, the label probes (first, second, etc.) can be hybridized to the corresponding amplifiers, and the cell can then be washed again prior to detection of the labels. As another example, the target probes can be hybridized to the targets, the cell can be washed, amplifiers and label probes can be added together and hybridized, and the cell can then be washed prior to detection. It will be evident that double-stranded nucleic acid target(s) are preferably denatured, e.g., by heat, prior to hybridization of the corresponding target probe(s) to the target(s).

In some embodiments, the cell is in suspension for all or most of the steps of the method, for ease of handling. However, the methods are also applicable to cells in solid tissue samples (e.g., tissue sections) and/or cells immobilized on a substrate (e.g., a slide or other surface). Thus, in one class of embodiments, the cell is in suspension in the sample comprising the cell, and/or the cell is in suspension during the hybridizing, capturing, and/or detecting steps. For example, the cell can be in suspension in the sample and during the hybridization, capture, optional washing, and detection steps. In other embodiments, the cell is in suspension in the sample comprising the cell, and the cell is fixed on a substrate during the hybridizing, capturing, and/or detecting steps. For example, the cell can be in suspension during the hybridization, capture, and optional washing steps and immobilized on a substrate during the detection step. In other embodiments, the sample comprises a tissue section.
3.1.2 Additional Nucleic Acid Detection Methods

RT-PCR. Methods of RT-PCR are known in the art and described, for example, in Dieffenbach (Ed) and Dveksler (Ed) (In: PCR Primer: A Laboratory Manual, Cold Spring Harbor Laboratories, NY, 1995).

Nanostring. Methods of Nanostring use labeled reporter molecules, referred to as labeled "nanoreporters," that are capable of binding individual target molecules. Through the nanoreporters' label codes, the binding of the nanoreporters to target molecules results in the identification of the target molecules. Methods of Nanostring are described in U.S. Patent No. 7,473,767.

Sequencing. The nucleic acid molecules can also be sequenced by any method known in the art, e.g., ensemble sequencing or single molecule sequencing. One conventional method to perform sequencing is by chain termination and gel separation, as described by Sanger et al., Proc Natl Acad Sci U S A, 74(12): 5463 67 (1977). Another conventional sequencing method involves chemical degradation of nucleic acid fragments. See, Maxam et al., Proc. Natl. Acad. Sci., 74: 560 564 (1977). Finally, methods have been developed based upon sequencing by hybridization. See, e.g., Drmanac, et al. (Nature Biotech., 16: 54 58, 1998).

qNPA™. The quantitative nuclease protection assay (qNPA™) is a method for measuring gene expression levels (mRNA) using gene-specific DNA oligonucleotides. S1 nuclease is used to degrade any non-hybridized nucleic acid, such as the non-hybridized portion of the targeted RNA, all of the non-targeted RNA, and excess DNA oligonucleotides. The resulting DNA oligos are a stoichiometrically representative library of the original RNA sample. The individual DNA probe oligonucleotides are present in the precise relative abundance as the RNA transcripts were in the original sample. Methods of qNPA™ are described at its manufacture's website: http://(www).htgmolecular.com.

QuantiGene® 2.0. The QuantiGene® 2.0 assay uses branched DNA (bDNA) technology. The bDNA assay is a sandwich nucleic acid hybridization method that uses bDNA molecules to amplify signal from captured target RNA. RNA is measured directly from the sample source in bDNA assays. Methods of QuantiGene® 2.0 assays are described at its manufacture's website: http://(www).panomics.com.
3.2 Protein Detection Methods

When the biological marker is a protein and/or a fragment and/or a variant thereof, several conventional methods for determining the level of a specific protein, and/or fragments and/or variants thereof, which are well-known to the skilled person, may be used. The level of the marker may for example be measured by using immunological assays, such as enzyme-linked immunosorbent assays (ELISA), thus providing a simple, reproducible and reliable method. Antibodies for use in such assays are available, and additional (polyclonal and monoclonal) antibodies may be developed using well-known standard techniques for developing antibodies. Other methods for measuring the level of the biological protein markers may furthermore include (immuno)histochemistry, Western blotting, flow-cytometry, RIA, competition assays, etc. and any combinations thereof.

3.2.1 Ligands and Antibodies

It will be apparent to the skilled artisan based on the disclosure herein that the present invention also extends to detection of a marker in a polypeptide, e.g., a target polypeptide encoded by a PHACTR1 mRNA. It is clear to a skilled artisan that detection of other target polypeptides such as SPP1, MLANA, PRAME, GDF15, CXCL10, S100A6, and S100B can be conducted in the same way as described below for PHACTR1.

Methods for detecting target polypeptides generally make use of a ligand or antibody that preferentially or specifically binds to the target polypeptide. As used herein the term "ligand" shall be taken in its broadest context to include any chemical compound, polynucleotide, peptide, protein, lipid, carbohydrate, small molecule, natural product, polymer, etc. that is capable of selectively binding, whether covalently or not, to one or more specific sites on an PHACTR1 polypeptide. The ligand may bind to its target via any means including hydrophobic interactions, hydrogen bonding, electrostatic interactions, van der Waals interactions, pi stacking, covalent bonding, or magnetic interactions amongst others.

As used herein, the term "antibody" refers to intact monoclonal or polyclonal antibodies, immunoglobulin (IgA, IgD, IgG, IgM, IgE) fractions, humanized antibodies, or recombinant single chain antibodies, as well as fragments thereof, such as, for example Fab, F(ab)2, and Fv fragments.
It will be apparent to the skilled artisan that the ligand and antibody against all the markers disclosed in the present invention can be prepared in the way as described below.

Antibodies are prepared by any of a variety of techniques known to those of ordinary skill in the art, and described, for example in, Harlow and Lane (In: Antibodies: A Laboratory Manual, Cold Spring Harbor Laboratory, 1988). In one such technique, an immunogen comprising the antigenic polypeptide is initially injected into any one of a wide variety of animals (e.g., mice, rats, rabbits, sheep, humans, dogs, pigs, chickens and goats). The immunogen is derived from a natural source, produced by recombinant expression means, or artificially generated, such as by chemical synthesis (e.g., BOC chemistry or FMOC chemistry).

A peptide, polypeptide or protein is joined to a carrier protein, such as bovine serum albumin or keyhole limpet hemocyanin. The immunogen and optionally a carrier for the protein is injected into the animal host, preferably according to a predetermined schedule incorporating one or more booster immunizations, and blood collected from said the animals periodically. Optionally, the immunogen is injected in the presence of an adjuvant, such as, for example Freund's complete or incomplete adjuvant, lysolecithin and dinitrophenol to enhance the subject's immune response to the immunogen. Monoclonal or polyclonal antibodies specific for the polypeptide are then purified from blood isolated from an animal by, for example, affinity chromatography using the polypeptide coupled to a suitable solid support.

Monoclonal antibodies specific for the antigenic polypeptide of interest are prepared, for example, using the technique of Kohler and Milstein, Eur. J. Immunol. 6:51 1-519, 1976, and improvements thereto. Briefly, these methods involve the preparation of immortal cell lines capable of producing antibodies having the desired specificity (i.e., reactivity with the polypeptide of interest). Such cell lines are produced, for example, from spleen cells obtained from an animal immunized as described supra. The spleen cells are immortalized by, for example, fusion with a myeloma cell fusion partner, preferably one that is syngenic with the immunized animal. A variety of fusion techniques are known in the art, for example, the spleen cells and myeloma cells are combined with a nonionic detergent or electrofused and then grown in a selective medium that supports the growth of hybrid cells, but not myeloma cells. A preferred selection technique uses HAT (hypoxanthine, aminopterin, and thymidine) selection. After a sufficient time, usually about 1 to 2 weeks, colonies of hybrids are observed. Single
colonies are selected and growth media in which the cells have been grown is tested for the presence of an antibody having binding activity against the polypeptide (immunogen). Hybridomas having high reactivity and specificity are preferred.

It is preferable that an immunogen used in the production of an antibody is one which is sufficiently antigenic to stimulate the production of antibodies that will bind to the immunogen and is preferably, a high titer antibody. In one embodiment, an immunogen is an entire protein.

In another embodiment, an immunogen consists of a peptide representing a fragment of a polypeptide, for example a region of a PHACTR1 polypeptide. Preferably an antibody raised to such an immunogen also recognizes the full-length protein from which the immunogen was derived, such as, for example, in its native state or having native conformation.

Alternatively, or in addition, an antibody raised against a peptide immunogen recognizes the full-length protein from which the immunogen was derived when the protein is denatured. By "denatured" is meant that conformational epitopes of the protein are disrupted under conditions that retain linear B cell epitopes of the protein. As will be known to a skilled artisan linear epitopes and conformational epitopes may overlap.

Alternatively, a monoclonal antibody capable of binding to a form of a PHACTR1 polypeptide or a fragment thereof is produced using a method such as, for example, a human B-cell hybridoma technique (Kozbar et al., Immunol. Today 4:72, 1983), a EBV-hybridoma technique to produce human monoclonal antibodies (Cole et al. "Monoclonal Antibodies in Cancer Therapy", 1985 Allen R. Bliss, Inc., pages 77-96), or screening of combinatorial antibody libraries (Huse et al., Science 246:1275, 1989).

Such an antibody is then particularly useful in detecting the presence of a marker of melanoma.

The methods described supra are also suitable for production of an antibody or antibody binding fragment as described herein according to any embodiment.

3.2.2 Detection Methods

In one embodiment, the method of the invention detects the presence of a marker in a polypeptide, aid marker being associated with melanoma.
An amount, level or presence of a polypeptide is determined using any of a variety of techniques known to the skilled artisan such as, for example, a technique selected from the group consisting of, immunohistochemistry, immunofluorescence, an immunoblot, a Western blot, a dot blot, an enzyme linked immunosorbent assay (ELISA), radioimmunoassay (RIA), enzyme immunoassay, fluorescence resonance energy transfer (FRET), matrix-assisted laser desorption/ionization time of flight (MALDI-TOF), electrospray ionization (ESI), mass spectrometry (including tandem mass spectrometry, e.g. LC MS/MS), biosensor technology, evanescent fiber-optics technology or protein chip technology.

In one example, an assay used to determine the amount or level of a protein is a semi-quantitative assay. In another example, an assay used to determine the amount or level of a protein in a quantitative assay.

Preferably, an amount of antibody or ligand bound to a marker of a melanoma in a PHACTR1 polypeptide is determined using an immunoassay. Preferably, using an assay selected from the group consisting of, immunohistochemistry, immunofluorescence, enzyme linked immunosorbent assay (ELISA), fluorescence linked immunosorbent assay (FLISA) Western blotting, RIA, a biosensor assay, a protein chip assay, a mass spectrometry assay, a fluorescence resonance energy transfer assay and an immunostaining assay (e.g. immunofluorescence).

4. Detection Kit

The present invention further relates to kits for performing the diagnostic methods as described above. The invention in particular relates to such diagnostic kits for identifying a subject with melanoma, comprising means for receiving one or more biological samples of said subject, and means for determining the level of the biological marker(s) in said biological sample of said subject. Thus a kit is provided which can be used as a reliable and easy diagnostic tool. One exemplary means for receiving the biological sample comprises a FFPE. The means for determining the presence of PHACTR1, SPP1, PRAME, GDF15, and CXCL10 genes in said biological sample of said subject may for example comprise one or more specific polynucleotide probes, antibodies, primers, etc. suitable for detecting the biological marker(s), identified according to the present invention. One exemplary means for determining the presence of the
marker genes is an RNAscope® assay kit equipped with mRNA markers which encode fragments of PHACTR1, SPPl, PRAME, GDF15, and CXCLIO genes, respectively. Another means for determining the presence of the marker genes is an RNAscope® assay kit equipped with mRNA markers which encode fragments of PHACTR1 and SPPl. The kits may further also comprise calibration means and instruction for use.

One general class of embodiments of the present invention provides a kit for detecting a nucleic acid target genes of PHACTR1, SPPl, PRAME, GDF15, and CXCLIO in an individual cell. One embodiment provides a kit for detecting a nucleic acid target genes of PHACTR1 and SPPl. The kits comprise one or more tissue pretreatment reagents, a target probe set, and a signal amplification system, packaged in one or more containers. The tissue pretreatment reagents comprise a citrate buffer and a protease. Tissue pretreatment procedure comprises heating the FFPE tissue sections in a citrate buffer followed by digestion with a protease. The purpose of these treatments is to allow for RNA access by the target probes through the disruption of formaldehyde cross-linking created in the sample preparation procedure. The target probe are designed to hybridize to each of the marker genes. In addition, probes against MLANA, S100A6 and S100B may be also included in the kit as positive controls, and a probe against the bacterial gene DapB may be included in the kit as negative control. The signal amplification system comprises, as described above in Section 3.1.1.4, pre-amplifier, amplifier, label probe and label. Each probe set contains a recognition sequence for a signal amplification system. The label probe refers to an entity that binds to the target molecule, directly or indirectly, and enables the target to be detected by a readout instrument. In one preferred embodiment, the label is Fast Red. Nucleic acid targets such as PHACTR1, SPPl, PRAME, GDF15, and CXCLIO can be detected by the formation of a precipitate following incubation with Fast Red.

Essentially all of the features noted for the embodiments in Sections 2.2 and 2.4 above apply to the kits as well, as relevant; for example, with respect to the identity of the markers, the expression level of the markers, the upregulation of the gene expression level or protein expression level of the markers, the patterns of the expression levels, and the type of gene expression analysis assays.
The features noted for the embodiments in Section 2.1.1.4 above apply to the kits as well, as relevant; for example, with respect to number of nucleic acid targets, configuration and number of the label and target probes, inclusion of preamplifiers and/or amplifiers, inclusion of blocking probes, inclusion of amplification reagents, type of nucleic acid target, location of various targets on a single molecule or on different molecules, type of labels, inclusion of optional blocking probes, and/or the like. The kit optionally also includes instructions for detecting the nucleic acid targets in the cell and/or identifying the cell as being of a specified type, one or more buffered solutions (e.g., diluent, hybridization buffer, and/or wash buffer), reference cell(s) comprising one or more of the nucleic acid targets, and/or the like.

One embodiment of the present invention provides a kit for detecting the expression level of PHACTR1 marker, either alone or in combination with one or more of SPPl, MLANA, PRAME, GDF15, CXCLIO, SI 10A, and S100B markers in a tissue sample, with all of the features noted for the embodiments above apply to this kit as well.

5. **Prognosis of Recurrence of Melanoma**

In addition to the initial detection of melanoma in a sample of a patient, the present invention can be used for prognosis of the likelihood of recurrence of melanoma in patients.

As disclosed above, the present invention uses a set of PHACTR1 and SPPl markers or a set of PHACTR1, SPPl, MLANA, PRAME, GDF15, and CXCLIO markers to determine the presence of melanoma. It is also discovered in the present invention that the same sets of marker genes can be used to predict the likelihood of recurrence of melanoma in patients.

In one embodiment, provided herein is a method providing a prognosis to a human subject who is at risk of developing melanoma recurrence, comprising the following steps:

(a) obtaining a tissue sample from said human subject, wherein said tissue sample comprises a plurality of melanocytes;

(b) measuring the level of expression of PHACTR1 marker and optionally the level of expression of one or more of SPPl, PRAME, GDF15, and CXCLIO markers; and

based on the gene expression levels of the PHACTR1 marker and the optionally one or more additional markers obtained for the tissue sample, providing a prognosis to the subject.
In another embodiment, level of expression of only PHACTR1 and SPP1 markers are used in the method for providing prognosis of melanoma.

All of the features noted for the embodiments described above in Sections 2.2 and 2.4 apply to the prognosis of recurrence of melanoma as well.

6. Determining the Stage of Melanoma Based on Sentinel Lymph Node

In addition to the diagnosis and prognosis of melanoma in a sample of a patient, the present invention can be used determine the stage of the melanoma patients. Currently, the best single prognostic factor for stage I-II disease is sentinel lymph node (SLN) status (Balch, CM. et al. (2004) "An Evidence-based Staging System for Cutaneous Melanoma". CA: A Cancer Journal for Clinicians 54, 131-149). Positive SLN status indicates presence of micrometastasis and portends future metastasis. However, the standard procedure of SLN assessment is based on histopathology, resulting in underestimating micrometastasis by 10-20%. RT-PCR has been proposed as a tool for molecular staging, but it uses markers not distinguishing between benign and malignant melanocytes and may not work well when there are only a few tumor cells present (Soikkeli, J. et al. (2007) "Systematic search for the best gene expression markers for melanoma micrometastasis detection" The Journal of Pathology 213, 180-9; Mocellin, S., Hoon, D.S.B., Pilati, P., Rossi, C.R. & Nitti, D. (2007) "Sentinel lymph node molecular ultrastaging in patients with melanoma: a systematic review and meta-analysis of prognosis" Journal of Clinical Oncology 25, 1588-95; Taback, B. et al. (2001) "The clinical utility of multimarker RT-PCR in the detection of occult metastasis in patients with melanoma" Recent Results in Cancer Research. 158:78-92).

As disclosed above, the present invention can detect positive staining in a few cells with and thus with high sensitivity. It can also differentiate melanoma with benign nevus with 100% accuracy. Further, the staining pattern of SPP1 can interrogate tumor morphology in both tumor and its microenvironment. Thus, it is discovered in one example of the invention, it has been surprisingly found that the biomarkers claimed in the present invention have the potential for prognosis of newly diagnosed melanoma patients by testing the patient's sentinel lymph node.

In one embodiment, provided herein is a method for determining the stage of melanoma in a human subject, comprising the following steps:
obtaining a tissue sample from said human subject, wherein said tissue sample comprises sentinel lymph node;

determining the level of expression of PHACTR1 marker and optionally the level of expression of one or more SPP1, PRAME, GDF15, and CXCL10 markers;

thereby determining the stage of melanoma as characterized by the presence of malignant melanoma cells in sentinel lymph node, based on the expression levels in said tissue sample.

In another embodiment, level of expression of only PHACTR1 and SPP1 markers are used in the method for determining the stage of melanoma.

All of the features noted for the embodiments described above in Sections 2.2 and 2.4 apply to the method for determining the stage of melanoma as well.

REFERENCES:


EXAMPLES

Example 1. Selection of PHACTR1, SPP1, and MLANA as Biomarkers for Detecting Melanoma

To identify the molecular events associated with melanoma progression and potential novel markers for early melanoma diagnosis, genome-wide gene expression microarrays were performed by several groups, and several datasets were made available through the Gene Expression Omnibus (GEO) database (http://www.ncbi.nlm.nih.gov/geo/). Although bioinformatic and statistical analysis had already been performed on the these datasets by the original authors, it was reasoned that combining results from multiple datasets, especially those from different microarray platforms, may lead to consistent, cross-validated candidate markers. Consequently, two different datasets comparing normal skin or benign nevi and malignant melanoma, GSE3189 (Affymetrix® GeneChip® HG-U133A, 22,000 genes) and GSE12391 (Agilent® Human Whole Genome Oligo Microarray, 41,000 genes) were selected, for candidate identification. GSE3189 contains data for 45 primary melanoma, 18 benign nevi and 7 normal skin specimens. GSE12391 contains a data series for 18 common nevi (CMN), 11 dysplastic nevi (DN), 8 radial (RGP) and 15 vertical (VGP) growth phase melanomas and 5 melanoma metastasis (MTS). These lesions represented different stages of melanoma progression according to a current model, which posits that melanoma initiates from dysplastic nevi, which then progresses to melanoma in situ or minimally invasive melanoma (RGP) to invasive melanoma (VGP) to metastatic melanoma (MTS) in a stepwise fashion.

The Limma linear models for microarrays were used to find genes differentially expressed between normal skin/benign nevi and melanomas. This algorithm uses empirical Bayes shrinkage of gene-wise residuals to ensure stable linear models when the number of samples is small, as is the case in these studies. PHACTR1 was the most and the second most strongly up-regulated gene in melanoma in GSE12391 and GSE3189, respectively (Table 1). The adjusted p values were after controlling for multiple hypothesis testing. The large difference in fold changes in these two datasets is likely due to the use of different microarray platforms and different data normalization schemes. Importantly, PHACTR1 demonstrated up-regulation in the early stage of RGP, whereas its expression in dysplastic nevi was similar to that in
common nevi (Fig. 4), suggesting its potential utility to differentiate between dysplastic nevi and melanomas. Interestingly, PHACTR1 was not selected for validation in either of the two original studies, probably due to its lower ranking by the different statistical tests used.

### Table 1. Consistent up-regulation of PHACTR1 in melanoma in two microarray datasets.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Dataset</th>
<th>Rank</th>
<th>log2(Fold Change)</th>
<th>t-statistic</th>
<th>Adjusted P</th>
</tr>
</thead>
<tbody>
<tr>
<td>PHACTR1</td>
<td>GSE12391</td>
<td>1</td>
<td>1.62</td>
<td>7.91</td>
<td>3.52E-06</td>
</tr>
<tr>
<td>PHACTR1</td>
<td>GSE3189</td>
<td>2</td>
<td>5.32</td>
<td>19.15</td>
<td>7.24E-26</td>
</tr>
</tbody>
</table>

Therefore, PHACTR1 was selected as the top candidate. In addition, SPP1 and four other genes were chosen for validation on the basis of their differential gene expression in these datasets and literature support. Finally, two widely used melanocyte-specific markers, MLANA and TYR were evaluated. The which MLANA and TYR genes are expressed both in normal melanocytes and melanoma cells, but not in other cell types in the skin.

### Example 2. Validation of PHACTR1, SPP1, and MLANA as Biomarkers for Detecting Melanoma

RNAscope® assays using gene-specific probes against these eight selected candidate genes were performed on a series of FFPE tissue microarrays (TMAs) obtained from commercial vendors. The bacterial gene DapB was used as negative control. These TMAs contained normal skin, nevi and melanomas of different stages. TMAs are efficient means to test many candidate genes in multiple samples.

In these TMA experiments, MLANA had higher and more consistent expression in benign nevi and melanomas than TYR. Thus, MLANA was chosen as the marker for melanocytes and melanoma cells. It also served as a positive control for RNA integrity, since these TMAs were constructed from archival tissues from various times and sources and had variable quality in RNA preservation. Together, 70% of the melanocytic nevi and melanomas on the TMAs had MLANA-staining intensity scores above 2+ on a semi-quantitative scale and were considered as having acceptable quality, resulting in 78 melanomas and 19 nevi in the final sample set.
Among the 6 candidate genes differentially expressed between nevi and melanomas, two genes, PHACTRl and SPPl, emerged as the strongest markers for melanoma and were detected at the early phase of melanoma (RGP). As shown in Table 2, 83% and 91% of the melanomas were positive for PHACTRl and SPPl, respectively, whereas none of the 19 nevi had detectable signals for either marker. In combination, 95% of the melanomas were positive for at least one marker: 84% were positive for both, 12% and 4% were positive for either SPPl or PHACTRl alone, respectively.

Table 2. Summary of three TMA studies combined.

<table>
<thead>
<tr>
<th>Sample Type</th>
<th>N</th>
<th>PHACTRl</th>
<th></th>
<th>SPPl</th>
<th></th>
<th>PHACTRl or SPPl</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Positive</td>
<td>%</td>
<td>Positive</td>
<td>%</td>
<td>Positive</td>
</tr>
<tr>
<td>Melanoma</td>
<td>78</td>
<td>65</td>
<td>83%</td>
<td>71</td>
<td>91%</td>
<td>74</td>
</tr>
<tr>
<td>Nevi</td>
<td>19</td>
<td>0</td>
<td>0%</td>
<td>0</td>
<td>0%</td>
<td>0</td>
</tr>
</tbody>
</table>

Examples of staining patterns of MLANA, PHACTRl and SPPl in a benign nevus and a melanoma are shown in Fig. 6. The benign nevus was completely negative for PHACTRl and SPPl, whereas the melanoma was positive for both.

Interestingly, it was noticed that in addition to the strong positive staining of SPPl in melanoma cells as shown in Fig. 7, in certain cases of melanoma, SPPl was detected primarily in individual cells scattered within the tumor region (Fig. 7). In a previous study using radioactive in situ hybridization, SPPl mRNA was detected specifically in tumor-associated macrophages (TAMs) in multiple tumor types including melanoma, especially at the interface between tumor and stroma. Therefore, it was demonstrated two patterns of SPPl expression: either predominantly in TAMs or in both TAMs and tumor cells. SPPl protein expression in melanoma cells was also observed using immunohistochemistry.

Example 3. Selection of Additional Biomarkers for Detecting Melanoma

Further improvements in RNAscope® Mela performance are highly desirable. First, MLANA was used as a positive control to help identify melanocytes (benign or malignant) and to assess quality of specimens. However, desmoplastic melanoma, representing 2-3% of all...
melanomas, is known to be negative for MLANA expression. A small fraction of other melanomas may also be negative for MLANA. Thus a negative MLANA staining result may be difficult to interpret. Second, PHACTR1 and SPP1 were negative in 5% of the melanomas, resulting in false negative classifications.

Therefore, it was sought to add additional markers to the 3-gene panel. First, two S100 genes, S100A6 and S100B, were added to MLANA to serve as positive control for assay procedure and sample quality control. S100A6 and S100B are known to be expressed in desmoplastic melanomas which are usually melan-A-negative. An example of RNAscope® detection of S100 in desmoplastic melanoma is shown in Fig. 8, where MLANA staining was weakly positive, but the pooled probes for S100A6 and S100B detected strong signals. In a TMA study, S100A6 and S100B staining patterns were highly correlated but some cases showed strong signals only in one of them. Therefore, the probes for MLANA, S100A6 and S100B were pooled together to serve as positive controls for melanocytes.

Next, 9 additional candidate genes (PRAME, SILV, CXCL9, CXCL10, SPRY4-IT1, GDF15, KRT15, PTN, and CFH) that are differentially expressed between melanoma and benign nevus based on literature review were selected. RNAscope® assays for the newly selected genes were developed and the 9 genes were tested for their diagnostic performance in a TMA of 100 samples (77 melanoma and 23 nevi). All staining were scored on a 0-4 semi-quantitative scale. Using a score cutoff of 3 or higher (3+), only 43% of the samples met this criterion using MLANA alone, but 70% of the samples met this criterion using the MLANA/S100A6/S100B probe pool, confirming the added sensitivity of S100 genes in the positive control. Among the 9 new and 2 existing genes (PHACTR1 and SPP1), PHACTR1 and SPP1 remain the best two single markers, followed by PRAME and GDF15 (see Fig. 9 for examples of RNAscope® staining), all showing positive staining in melanomas and negative staining in nevi, with no false positives (Table 3). The two chemokines CXCL9 and CXCL10 also showed 100% specificity but their sensitivity was low (Table 3). However, these two genes are stromal markers and expressed in tumor-associated endothelial cells (see Fig. 9 for an example of RNAscope® staining). Since CXCL10 showed somewhat higher signals and sensitivity than CXCL9, it was chosen for further evaluation.
Example 4. Evaluation of PHACTR1, SPPl, MLANA, PRAME, GDF15, CXCLIO, S100A6 and S100B as Biomarkers for Detecting Melanoma

Potential combinations of PHACTR1, SPPl, MLANA, PRAME, GDF15, CXCLIO, S100A6 and S100B genes to increase sensitivity while maintaining specificity at 100% (Table 3). Since both CXCLIO and SPPl are expressed in the tumor-associated stroma, although in different cell types, it was sought to combine these two markers. Although combining SPPl with CXCLIO did not lead to further increase in sensitivity compared to SPPl alone, the combined macrophage- and endothelial-staining cells makes the assay more robust. Compared to the original proposed panel consisting of PHACTR1 and SPPl, adding PRAME, GDF15 and CXCLIO resulted in an increase in sensitivity from 95% to 98%, while maintaining 100% specificity. Interestingly, logistic regression with model selection resulted in the minimal combination of PRAME and SPPl, which had the same level of performance as the 5-gene panel. The combination of 5 genes of PHACTR1, SPPl, PRAME, GDF15 and CXCLIO demonstrated the potential of a reduced model for melanoma detection.

Table 3. Classification performance of individual markers and their combinations.

<table>
<thead>
<tr>
<th>Pos. control=3+, nevi=8, mm=62</th>
<th>Sensitivity</th>
<th>Specificity</th>
<th>PPV</th>
<th>NPV</th>
</tr>
</thead>
<tbody>
<tr>
<td>SILV</td>
<td>0.87</td>
<td>0.25</td>
<td>0.90</td>
<td>0.20</td>
</tr>
<tr>
<td>CXCL9</td>
<td>0.21</td>
<td>1.00</td>
<td>1.00</td>
<td>0.14</td>
</tr>
<tr>
<td>SPRY4.1T1</td>
<td>0.30</td>
<td>1.00</td>
<td>1.00</td>
<td>0.16</td>
</tr>
<tr>
<td>CXCLIO</td>
<td>0.36</td>
<td>1.00</td>
<td>1.00</td>
<td>0.17</td>
</tr>
<tr>
<td>SPPl</td>
<td>0.95</td>
<td>1.00</td>
<td>1.00</td>
<td>0.73</td>
</tr>
<tr>
<td>PHACTR1</td>
<td>0.79</td>
<td>1.00</td>
<td>1.00</td>
<td>0.73</td>
</tr>
<tr>
<td>PRAME</td>
<td>0.72</td>
<td>1.00</td>
<td>1.00</td>
<td>0.32</td>
</tr>
<tr>
<td>GDF15</td>
<td>0.64</td>
<td>1.00</td>
<td>1.00</td>
<td>0.32</td>
</tr>
<tr>
<td>KRT15</td>
<td>0.95</td>
<td>0.63</td>
<td>0.95</td>
<td>0.63</td>
</tr>
<tr>
<td>CFH</td>
<td>0.39</td>
<td>1.00</td>
<td>1.00</td>
<td>0.18</td>
</tr>
<tr>
<td>PTN</td>
<td>0.97</td>
<td>0.75</td>
<td>0.97</td>
<td>0.75</td>
</tr>
<tr>
<td>PHACTR1.PRAME</td>
<td>0.89</td>
<td>1.00</td>
<td>1.00</td>
<td>0.53</td>
</tr>
<tr>
<td>PHACTR1.GDF15</td>
<td>0.89</td>
<td>1.00</td>
<td>1.00</td>
<td>0.53</td>
</tr>
<tr>
<td>PHACTR1.GDF15.PRAME</td>
<td>0.93</td>
<td>1.00</td>
<td>1.00</td>
<td>0.67</td>
</tr>
<tr>
<td>SPPl.CXCLIO</td>
<td>0.95</td>
<td>1.00</td>
<td>1.00</td>
<td>0.73</td>
</tr>
<tr>
<td>PHACTR1.GDF15.PRAME.SPP1.CXCL10</td>
<td>0.98</td>
<td>1.00</td>
<td>1.00</td>
<td>0.89</td>
</tr>
<tr>
<td>PHACTR1.SPP1</td>
<td>0.95</td>
<td>1.00</td>
<td>1.00</td>
<td>0.73</td>
</tr>
<tr>
<td>PRAME.SPP1</td>
<td>0.98</td>
<td>1.00</td>
<td>1.00</td>
<td>0.89</td>
</tr>
</tbody>
</table>
We also performed an analysis between the expression levels of 5 genes of PHACTR1, SPPI, PRAME, GDF15 and CXCLIO and their relations to the diagnosis of melanoma. We tested for associations between the copy number of mRNA of each above gene and the presence of melanoma. We found that in patients with melanoma, the copy number of mRNA of PHACTR1, PRAME, GDF15 and CXCLIO genes could range from 1 to over 50. The copy number of mRNA of SPPI gene can be even higher, ranging from 1 to over 100. We determined that 1 copy of mRNA can serve as a reliable threshold for determining the presence of melanoma in a tissue sample, meaning that if there is a single copy of mRNA of any of the 5 genes of PHACTR1, SPPI, PRAME, GDF15 and CXCLIO, melanoma can be reliably predicted. We also found that if the threshold is raised to 3 copies of mRNA of any one of the 5 genes, the presence of melanoma can be predicted in certain.

Although the present invention has been described in detail with reference to the specific features, it will be apparent to those skilled in the art that this description is only for a preferred embodiment and does not limit the scope of the present invention. Various modifications may be made without deviating from the spirit and scope of what is provided herein.
WHAT IS CLAIMED IS:

1. A method for diagnosing melanoma in a human subject suspected of melanoma, comprising the following steps:
   (a) obtaining a tissue sample from said human subject, wherein said tissue sample comprises a plurality of melanocytes;
   (b) determining the level of expression of at least phosphatase and actin regulator 1 (PHACTR1) gene, or fragments thereof; and
   (c) optionally determining the level of expression of one or more additional markers of melanoma, or fragments thereof; thereby diagnosing the presence of melanoma based on the expression levels in said tissue sample.

2. The method of claim 1, wherein said determining the level of expression comprises determining the level of expression of a set of genes comprising PHACTR1 and one or more of genes selected from the group consisting of secreted integrin-binding phosphoprotein (SPP1), preferentially expressed antigen in melanoma (PRAME), growth differentiation factor 15 (GDF15), and chemokine C-X-C motif ligand 10 (CXCL10) genes.

3. The method of claim 2, wherein said determining the level of expression comprises determining the level of PHACTR1 and SPP1 genes.

4. The method of any one of claims 1-3, wherein said determining the level of expression comprises detecting the expression level of mRNA expressed from said genes.

5. The method of any one of claims 1-3, wherein said determining the level of expression comprises detecting the expression level of polypeptide encoded by said genes.

6. The method of claim 2 or 3, wherein diagnosing the presence of melanoma is based on a pattern of expression levels of the set of genes.
7. The method of claim 6, wherein the levels of expression of the set of genes are determined individually.

8. The method of claim 6, wherein the levels of expression of the set of genes are pooled and determined altogether.

9. The method of claim 4, wherein the upregulation of the mRNA expression of said genes is indicative of melanoma.

10. The method of claim 5, wherein the upregulation of the polypeptide expression of said genes is indicative of melanoma.

11. The method of claim 2, wherein the presence of PHACTR1 and one or more of genes selected from the group consisting of SPPl, PRAME, GDF15, and CXCLIO genes is indicative of the presence of melanoma but the absence of benign nevus in said human subject.

12. The method of claim 3, wherein the presence of PHACTR1 and SPPl genes is indicative of the presence of melanoma but the absence of benign nevus in said human subject.

13. The method of claim 4, wherein said determining the level of expression comprises detection of mRNA using a gene expression analysis assay which measures mRNA in solution.

14. The method of claim 13, wherein said determining the level of expression is performed \textit{in situ}.

15. The method of claim 13, wherein the gene expression analysis assay comprises RNAscope®, RT-PCR, Nanostring, QuantiGene® 2.0, qNPA™, and microarray.

16. The method of any one of claims 1-3, wherein said tissue sample is skin.

17. A method for providing a prognosis to a human subject who is at risk of developing melanoma recurrence, comprising the following steps:
(a) obtaining a tissue sample from said human subject, wherein said tissue sample comprises a plurality of melanocytes;

(b) determining the level of expression of at least PHACTR1 gene, or fragments thereof;

(c) optionally determining the level of expression of one or more additional markers of melanoma, or fragments thereof; and

(d) based on the level of expression of PHACTR1 and the optionally one or more additional markers obtained for the tissue sample, providing a prognosis to the subject.

18. The method of claim 17, wherein said determining the level of expression comprises determining the level of expression of a set of genes comprising PHACTR1 and one or more of genes selected from the group consisting of SPP1, PRAME, GDF15, and CXCL10 genes.

19. The method of claim 18, wherein said determining the level of expression comprises determining the level of PHACTR1 and SPP1 genes.

20. The method of any one of claims 17-19, wherein said determining the level of expression of the genes comprises detecting the expression level of mRNA expressed from said genes.

21. The method of any one of claims 17-19, wherein said determining the level of expression of the genes comprises detecting the expression level of polypeptide encoded by said genes.

22. The method of claim 18 or 19, wherein said providing a prognosis to the subject is based on a pattern of expression levels of the set of genes.

23. The method of claim 22, wherein the levels of expression of the set of genes are determined individually.

24. The method of claim 22, wherein the levels of expression of the set of genes are pooled and determined altogether.
25. The method of claim 20, wherein the upregulation of the mRNA expression of said genes is indicative of recurrence of melanoma.

26. The method of claim 21, wherein the upregulation of the polypeptide expression of said genes is indicative of recurrence of melanoma.

27. The method of claim 20, wherein said determining the level of expression comprises detection of mRNA using a gene expression analysis assay which measures mRNA in solution.

28. The method of claim 27, wherein said determining the level of expression is performed in situ.

29. The method of claim 27, wherein the gene expression analysis assay comprises: RNAscope®, RT-PCR, Nanostring, QuantiGene® 2.0, qNPA™, and microarray.

30. A method for determining the stage of melanoma in a human subject, comprising the following steps:

   (a) obtaining a tissue sample from said human subject, wherein said tissue sample comprises sentinel lymph node;

   (b) determining the level of expression of at least PHACTR1 gene, or fragments thereof; and

   (c) optionally determining the level of expression of one or more additional markers of melanoma, or fragments thereof;

   thereby determining the stage of melanoma as characterized by the presence of malignant melanoma cells in sentinel lymph node, based on the expression levels in said tissue sample.

31. The method of claim 30, wherein said determining the level of expression comprises determining the level of expression of a set of genes comprising PHACTR1 and one or more of genes selected from the group consisting of SPP1, PRAME, GDF15, and CXCL10 genes.
32. The method of claim 30, wherein said determining the level of expression comprises determining the level of expression of PHACTR1 and SPP1 genes.

33. The method of any one of claims 30-32, wherein said determining the level of expression comprises detecting the expression level of mRNA expressed from said genes.

34. The method of any one of claims 30-32, wherein said determining the level of expression comprises detecting the expression level of polypeptide encoded by said genes.

35. The method of claim 31 or 32, wherein said determining the stage of melanoma is based on a pattern of expression levels of the set of genes.

36. The method of claim 35, wherein the levels of expression of the set of genes are determined individually.

37. The method of claim 35, wherein the levels of expression of the set of genes are pooled and determined altogether.

38. The method of claim 33, wherein the upregulation of the mRNA expression of said genes is indicative of risk of recurrence of melanoma.

39. The method of claim 34, wherein the upregulation of the polypeptide expression of said genes is indicative risk of recurrence of melanoma.

40. The method of claim 32, wherein said determining the level of expression comprises detection of mRNA using a gene expression analysis assay which measures mRNA in solution.

41. The method of claim 40, wherein said determining the level of expression is performed in situ.

42. The method of claim 40, wherein the gene expression analysis assay comprises RNAscope®, RT-PCR, Nanostring, QuantiGene® 2.0, qNPA™, and microarray.
43. A diagnostic kit, comprising at least one substance for detection of the level of expression of PHACTR1 gene, either alone or in combination with the detection of one or more genes selected from the group consisting of SPPL, PRAME, GDF15, and CXCLIO, for the diagnosis or prognosis of melanoma.

44. The diagnostic kit of claim 43, wherein the kit comprises a gene expression analysis means which measures mRNA in solution.

45. The method of claim 44, wherein said determining the level of expression is performed in situ.

46. The diagnostic kit of claim 44, wherein said gene expression analysis assay comprises RNAscope®, RT-PCR, Nanostring, QuantiGene® 2.0, qNPA™, and microarray.

47. The diagnostic kit of claim 43, wherein the kit comprises a protein expression analysis means which measures the expression level of polypeptide encoded by said genes.
Fig. 2
Fig. 4

A

PHACTR1: GSE12391

P = 5.6e-07

Dab2

Log2(Ratio)

MTS

vGP

RGP

DN

CMN

B

PHACTR1: GSE3189

P = 3.8e-11

Melanoma

Nevus

Normal

Log2(Expression)
Fig. 6