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(71) **Demandeur/Applicant:**
OXFORD BIODYNAMICS LIMITED, GB

(72) **Inventeurs/Inventors:**
AKOULITCHEV, ALEXANDRE, GB;
RAMADASS, AROUL SELVAM, GB;
HUNTER, EWAN, GB;
SALTER, MATTHEW, GB

(74) **Agent:** DEETH WILLIAMS WALL LLP

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A process for analysing chromosome regions and interactions relating to ALS and Huntington's disease.

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(71) Applicant: **OXFORD BIODYNAMICS LIMITED**
[GB/GB]; 26 Beaumont Street, Oxford Oxfordshire OX1 2NP (GB).

(72) Inventors: **AKOULITCHEV, Alexandre**; c/o Oxford BioDynamics Limited, 26 Beaumont Street, Oxford Oxfordshire OX1 2NP (GB). **RAMADASS, Aroul, Selvam**; c/o Oxford BioDynamics Limited, 26 Beaumont Street, Oxford Oxfordshire OX1 2NP (GB). **HUNTER, Ewan**; c/o Oxford BioDynamics Limited, 26 Beaumont Street, Oxford Oxfordshire OX1 2NP (GB). **SALTER, Matthew**; c/o Oxford BioDynamics Limited, 26 Beaumont Street, Oxford Oxfordshire OX1 2NP (GB).

(74) Agent: **AVIDITY IP**; Broers Building, Hauser Forum, 21 J J Thomson Ave, Cambridge Cambridgeshire CB3 0FA (GB).

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(54) Title: BIOMARKER

(57) Abstract: A process for analysing chromosome regions and interactions relating to ALS and Huntington's disease.

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BIOMARKER**Field of the Invention**

The invention relates to detecting chromosome interactions.

Background of the Invention

5 Biomarkers allow disease characteristics to be identified. The present commonly used biomarkers include RNA expression patterns and protein markers.

Summary of the Invention

Specific Chromosome Conformation Signatures (CCSs) at loci either exist or are absent due to the regulatory epigenetic control settings associated with pathology or treatment. CCSs have mild off-rates
10 and when representing a particular phenotype or pathology, they will only change with a physiologically signalled transition to a new phenotype, or as a result of external intervention. In addition, the measurement of these events is binary, and so this read-out is in stark contrast to the continuum readout of varying levels of DNA methylation, histone modifications and most of the non-coding RNAs. The continuum read-out used for most molecular biomarkers to date offers a challenge to data analysis, in
15 that the magnitude of change for particular biomarkers varies greatly from patient to patient, which causes problems for classification statistics when they are used to stratify cohorts of patients. These classification statistics are better-suited to using biomarkers that are absent of magnitude and offer just a “yes or no” binary score of phenotypic differences - signifying that chromosome conformation (*EpiSwitch™*) biomarkers are an excellent resource for potential diagnostic, prognostic and predictive
20 biomarkers.

The inventors have identified regions of the genome where chromosomal interactions are relevant to amyotrophic lateral sclerosis (ALS) or Huntington’s disease using an approach which allows identification of subgroups in a population. Accordingly, the invention provides a process for detecting a chromosome
25 state which represents a subgroup in a population comprising determining whether a chromosome interaction is present or absent within a defined disease-associated region of the genome, wherein said disease is ALS or Huntington’s disease. The chromosome interaction may optionally have been identified, or be identifiable (or derivable), by a method of determining which chromosomal interactions are relevant to a chromosome state corresponding to an ALS or Huntington’s disease subgroup of the population,
30 comprising contacting a first set of nucleic acids from subgroups with different states of the chromosome with a second set of index nucleic acids, and allowing complementary sequences to hybridise, wherein the nucleic acids in the first and second sets of nucleic acids represent a ligated product comprising sequences from both the chromosome regions that have come together in chromosomal interactions, and wherein the pattern of hybridisation between the first and second set of nucleic acids allows a

determination of which chromosomal interactions are specific to a ALS or Huntington's disease subgroup. The ALS or Huntington's disease subgroup may relate to diagnosis (presence of ALS or Huntington's disease) or prognosis (for example rate of progress of ALS or Huntington's disease). Any of the specific relevant chromosome interactions (markers) described herein may be used as the basis of the invention,
5 including combinations of markers.

The invention provides a process for detecting a chromosome state which represents a disease subgroup in a population comprising determining whether a chromosome interaction relating to that chromosome state is present or absent within a defined region of the genome, wherein said disease subgroup is an
10 amyotrophic lateral sclerosis (ALS) subgroup; and

- wherein said chromosome interaction has optionally been identified by a method of determining which chromosomal interactions are relevant to a chromosome state corresponding to an ALS subgroup of the population, comprising contacting a first set of nucleic acids from subgroups with different states of the chromosome with a second set of index nucleic acids, and allowing complementary sequences to
15 hybridise, wherein the nucleic acids in the first and second sets of nucleic acids represent a ligated product comprising sequences from both the chromosome regions that have come together in chromosomal interactions, and wherein the pattern of hybridisation between the first and second set of nucleic acids allows a determination of which chromosomal interactions are specific to an ALS subgroup; and
- wherein the chromosome interaction:
 - 20 (i) is present in any one of the regions or genes listed in Table 1 or 5; and/or
 - (ii) corresponds to any one of the chromosome interactions represented by any probe shown in Table 1 or 5, and/or
 - (iii) corresponds to any one of the chromosome interactions shown in Table 10 or 11, and/or
 - (iv) is present in a 4,000 base region which comprises or which flanks (i), (ii) or (iii).

25 The invention also provides a process for detecting a chromosome state which represents a disease subgroup in a population comprising determining whether a chromosome interaction relating to that chromosome state is present or absent within a defined region of the genome, wherein said disease subgroup is a Huntington's disease subgroup; and

30 - wherein said chromosome interaction has optionally been identified by a method of determining which chromosomal interactions are relevant to a chromosome state corresponding to an Huntington's disease subgroup of the population, comprising contacting a first set of nucleic acids from subgroups with different states of the chromosome with a second set of index nucleic acids, and allowing complementary sequences to hybridise, wherein the nucleic acids in the first and second sets of nucleic acids represent a
35 ligated product comprising sequences from both the chromosome regions that have come together in chromosomal interactions, and wherein the pattern of hybridisation between the first and second set of

nucleic acids allows a determination of which chromosomal interactions are specific to an Huntington's disease subgroup; and

- wherein the chromosome interaction:

(i) is present in any one of the regions or genes listed in Table 12; and/or

5 (ii) corresponds to any one of the chromosome interactions represented by any probe shown in Table 12, and/or

(iii) corresponds to any one of the chromosome interactions represented in Table 12, and/or

(iii) is present in a 4,000 base region which comprises or which flanks (i), (ii) or (iii).

10 **Brief Description of the Drawings**

Figure 1 shows how the chromosome interaction can be detected.

Figure 2 shows a visual overview of the genomic region investigated in the Huntington's disease work. A

~225 kb region on chromosome 4 spanning the HTT locus was investigated. The Anchor point ("Anchor"

15 in track 4) was defined as a ~42 kb region spanning the CAG repeat tract in exon 1 of HTT (purple arrow

at the top of the figure). We defined five Zones (Zones 1-5 in track 4) based on overlap with *EpiSwitch* sites (track 3), SNPs related to HD (track 5) or other diseases (track 6), and observed methylation and acetylation (H3K4me3, H3K36me3 and H3K27Ac) differences between HC and HD (tracks 7 through 12).

Figure 3 shows that in six out of seven individual HD-Sym samples, the presence of at least one of the

three conditional interactions (I5, I6 and I7) was observed. I5, the interaction spanning the region that

20 contains the rs362331 SNP, was observed in the greatest number of samples (6/7).

Figure 4 provides a model for epigenetic changes driving the symptomatic progression of Huntington's disease. It shows an overview of the chromosomal conformation changes associated with the progression of HD. As patients progress from presymptomatic stages to symptomatic diseases, discrete, measurable and discriminating changes in the genomic architecture at the HTT locus are observed.

25 Figure 5 shows a summary of all the Huntington's disease interactions which were identified.

Detailed Description of the Invention

The Process of the Invention

The process of the invention comprises a typing system for detecting chromosome interactions relevant

to ALS or Huntington's disease. This typing may be performed using the *EpiSwitch*™ system mentioned

30 herein which is based on cross-linking regions of chromosome which have come together in the chromosome interaction, subjecting the chromosomal DNA to cleavage and then ligating the nucleic acids present in the cross-linked entity to derive a ligated nucleic acid with sequence from both the regions

which formed the chromosomal interaction. Detection of this ligated nucleic acid allows determination of the presence or absence of a particular chromosome interaction.

The chromosomal interactions may be identified using the above described method in which populations of first and second nucleic acids are used. These nucleic acids can also be generated using *EpiSwitch*™

5 technology.

The Epigenetic Interactions Relevant to the Invention

As used herein, the term 'epigenetic' and 'chromosome' interactions typically refers to interactions between distal regions of a chromosome, said interactions being dynamic and altering, forming or breaking depending upon the status of the region of the chromosome.

10 In particular processes of the invention chromosome interactions are detected by first generating a ligated nucleic acid that comprises sequence from both regions of the chromosomes that are part of the interactions. In such processes the regions can be cross-linked by any suitable means. In a preferred embodiment, the interactions are cross-linked using formaldehyde, but may also be cross-linked by any aldehyde, or D-Biotinoyl-e- aminocaproic acid-N-hydroxysuccinimide ester or Digoxigenin-3-O-15 methylcarbonyl-e-aminocaproic acid-N-hydroxysuccinimide ester. Para-formaldehyde can cross link DNA chains which are 4 Angstroms apart. Preferably the chromosome interactions are on the same chromosome and optionally 2 to 10 Angstroms apart.

20 The chromosome interaction may reflect the status of the region of the chromosome, for example, if it is being transcribed or repressed in response to change of the physiological conditions. Chromosome interactions which are specific to subgroups as defined herein have been found to be stable, thus providing a reliable means of measuring the differences between the two subgroups.

25 In addition, chromosome interactions specific to a characteristic (such as a disease condition) will normally occur early in a biological process, for example compared to other epigenetic markers such as methylation or changes to binding of histone proteins. Thus the process of the invention is able to detect early stages of a biological process. This allows early intervention (for example treatment) which may as a consequence be more effective. Furthermore there is little variation in the relevant chromosome interactions between individuals within the same subgroup. Detecting chromosome interactions is highly informative with up to 50 different possible interactions per gene, and so processes of the invention can interrogate 500,000 different interactions.

30 *Preferred Marker Sets*

Herein the term 'marker' or 'biomarker' refers to a specific chromosome interaction which can be detected (typed) in the invention. Specific markers are disclosed herein, any of which may be used in the invention. Further sets of markers may be used, for example in the combinations or numbers disclosed

herein. The markers disclosed in the tables herein are preferred. These may be typed by any suitable method, for example the PCR or probe based methods disclosed herein, including a qPCR method. The markers are defined herein by location or by probe and/or primer sequences.

Location and Causes of Epigenetic Interactions

- 5 Epigenetic chromosomal interactions may overlap and include the regions of chromosomes shown to encode relevant or undescribed genes, but equally may be in intergenic regions. It should further be noted that the inventors have discovered that epigenetic interactions in all regions are equally important in determining the status of the chromosomal locus. These interactions are not necessarily in the coding region of a particular gene located at the locus and may be in intergenic regions.
- 10 The chromosome interactions which are detected in the invention could be caused by changes to the underlying DNA sequence, by environmental factors, DNA methylation, non-coding antisense RNA transcripts, non-mutagenic carcinogens, histone modifications, chromatin remodelling and specific local DNA interactions. The changes which lead to the chromosome interactions may be caused by changes to the underlying nucleic acid sequence, which themselves do not directly affect a gene product or the
- 15 mode of gene expression. Such changes may be for example, SNPs within and/or outside of the genes, gene fusions and/or deletions of intergenic DNA, microRNA, and non-coding RNA. For example, it is known that roughly 20% of SNPs are in non-coding regions, and therefore the process as described is also informative in non-coding situation. In one embodiment the regions of the chromosome which come together to form the interaction are less than 5 kb, 3 kb, 1 kb, 500 base pairs or 200 base pairs
- 20 apart on the same chromosome.

The chromosome interaction which is detected is preferably within any of the genes mentioned in Table 1 or 5. However it may also be upstream or downstream of the gene, for example up to 50,000, up to 30,000, up to 20,000, up to 10,000 or up to 5000 bases upstream or downstream from the gene or from the coding sequence.

25 *Subgroups, Diagnosis and Personalised Treatment*

The aim of the present invention is to permit detection of chromosome interactions relevant to an ALS or Huntington's disease subgroup. Therefore the process may or may not be used for diagnosis of ALS or Huntington's disease. The process may or may not be used for prognosis of ALS or Huntington's disease.

- When the process is used for diagnosis of ALS the typing of markers relating to Table 1 are preferred (i.e. 30 the specific disclosed markers and those in the genes, regions and flanking areas disclosed in Table 1). In one embodiment relating to diagnosis only markers relating to Table 1 are typed and no other markers. In other embodiments relating to diagnosis at least 1, 2, 3, 4, 5 or more markers relating to (for example as represented by the listed probe sequences) Table 5 are not typed.

When the process is used for prognosis the typing of markers relating to Table 5 are preferred (i.e. the specific disclosed markers and those in the genes, regions and flanking areas disclosed in Table 5). In one embodiment relating to prognosis only markers relating to Table 5 are typed and no other markers. In other embodiments relating to prognosis at least 1, 2, 3, 4, 5 or more markers relating to (for example as 5 represented by the listed probe sequences) Table 1 are not typed.

Typically, 'prognosis' relates to progression of ALS, and allows individuals to be divided into rate of progression subgroups. Progression may be measured using an ALS-FRS-R score (ALS functional rating scale), and individuals may be classed as being above or below a certain value, for example above or below a 0.5 point decline per 30 days. This allows a predictive prognosis to be performed.

10 As used herein, a "subgroup" preferably refers to a population subgroup (a subgroup in a population), more preferably a subgroup in the population of a particular animal such as a particular eukaryote, or mammal (e.g. human, non-human, non-human primate, or rodent e.g. mouse or rat). Most preferably, a "subgroup" refers to a subgroup in the human population.

The invention includes detecting and treating particular subgroups in a population. The inventors have 15 discovered that chromosome interactions differ between subsets (for example at least two subsets) in a given population. Identifying these differences will allow physicians to categorize their patients as a part of one subset of the population as described in the process. The invention therefore provides physicians with a process of personalizing medicine for the patient based on their epigenetic chromosome interactions.

20 In one embodiment subgroups (for example relating to prognosis) are defined by the ALS functional rating scale, for example as described in Cedarbaum, J.M., Stambler, N., Malta, E., Fuller, C., Hilt, D., Thurmond, B. and Nakanishi, A. (1999) The ALSFRS-R: a revised ALS functional rating scale that incorporates assessments of respiratory function. BDNF ALS Study Group (Phase III), *Journal of the Neurological Sciences*. 169(1-2), 13-21.

25 In one embodiment subgroups (for example relating to prognosis) are defined by forced vital capacity (FVC), for example as described in Talakad, N.S., Pradhan, C., Nalini, A., Thennarasu, K. and Raju T.R. (2009) Assessment of Pulmonary Function in Amyotrophic Lateral Sclerosis, *Indian J Chest Dis Allied Sci*. 51(2):87-91.

Generating Ligated Nucleic Acids

30 Certain embodiments of the invention utilise ligated nucleic acids, in particular ligated DNA. These comprise sequences from both of the regions that come together in a chromosome interaction and therefore provide information about the interaction. The EpiSwitch™ method described herein uses generation of such ligated nucleic acids to detect chromosome interactions.

Thus a process of the invention may comprise a step of generating ligated nucleic acids (e.g. DNA) by the following steps (including a method comprising these steps):

(i) cross-linking of epigenetic chromosomal interactions present at the chromosomal locus, preferably *in vitro*;

5 (ii) optionally isolating the cross-linked DNA from said chromosomal locus;

(iii) subjecting said cross-linked DNA to cutting, for example by restriction digestion with an enzyme that cuts it at least once (in particular an enzyme that cuts at least once within said chromosomal locus);

(iv) ligating said cross-linked cleaved DNA ends (in particular to form DNA loops); and

(v) optionally identifying the presence of said ligated DNA and/or said DNA loops, in particular using 10 techniques such as PCR (polymerase chain reaction), to identify the presence of a specific chromosomal interaction.

These steps may be carried out to detect the chromosome interactions for any embodiment mentioned herein, such as for determining whether the individual is part of an ALS or Huntington's disease subgroup.

The steps may also be carried out to generate the first and/or second set of nucleic acids mentioned 15 herein.

PCR (polymerase chain reaction) may be used to detect or identify the ligated nucleic acid, for example the size of the PCR product produced may be indicative of the specific chromosome interaction which is present, and may therefore be used to identify the status of the locus. In preferred embodiments at least 1, 2, 3, 4, 5, 6, 7 or 8 primers or primer pairs as shown in Table 2 or 7 are used in the PCR reaction. In other

20 preferred embodiments at least 1, 2, 3, 4, 5, 6, 7 or 8 primers or primer pairs as shown in any other table are used in the PCR reaction. The skilled person will be aware of numerous restriction enzymes which can be used to cut the DNA within the chromosomal locus of interest. It will be apparent that the particular enzyme used will depend upon the locus studied and the sequence of the DNA located therein. A non-limiting example of a restriction enzyme which can be used to cut the DNA as described in the present 25 invention is TaqI.

Embodiments such as EpiSwitch™ Technology

The EpiSwitch™ Technology also relates to the use of microarray EpiSwitch™ marker data in the detection of epigenetic chromosome conformation signatures specific for phenotypes. Embodiments such as EpiSwitch™ which utilise ligated nucleic acids in the manner described herein have several advantages.

30 They have a low level of stochastic noise, for example because the nucleic acid sequences from the first set of nucleic acids of the present invention either hybridise or fail to hybridise with the second set of nucleic acids. This provides a binary result permitting a relatively simple way to measure a complex

mechanism at the epigenetic level. EpiSwitch™ technology also has fast processing time and low cost. In one embodiment the processing time is 3 hours to 6 hours.

Samples and Sample Treatment

The process of the invention will normally be carried out on a sample. The sample will normally contain

5 DNA from the individual. It will normally contain cells. In one embodiment a sample is obtained by minimally invasive means, and may for example be a blood sample. DNA may be extracted and cut up with a standard restriction enzyme. This can pre-determine which chromosome conformations are retained and will be detected with the EpiSwitch™ platforms. Due to the synchronisation of chromosome interactions between tissues and blood, including horizontal transfer, a blood sample can be used to

10 detect the chromosome interactions in tissues, such as tissues relevant to disease. For certain conditions, such as cancer, genetic noise due to mutations can affect the chromosome interaction 'signal' in the relevant tissues and therefore using blood is advantageous.

Properties of Nucleic Acids of the Invention

15 The invention relates to certain nucleic acids, such as the ligated nucleic acids which are described herein as being used or generated in the process of the invention. These may be the same as, or have any of the properties of, the first and second nucleic acids mentioned herein. The nucleic acids of the invention typically comprise two portions each comprising sequence from one of the two regions of the chromosome which come together in the chromosome interaction. Typically each portion is at least 8, 10,

20 15, 20, 30 or 40 nucleotides in length, for example 10 to 40 nucleotides in length. Preferred nucleic acids comprise sequence from any of the genes mentioned in any of the tables. Typically preferred nucleic acids comprise the specific probe sequences mentioned in Table 1 or 5; or fragments and/or homologues of such sequences. Other preferred nucleic acids comprise the specific probe sequences mentioned in Table 10, 11 or 12; or fragments and/or homologues of such sequences. Preferably the nucleic acids are DNA.

25 It is understood that where a specific sequence is provided the invention may use the complementary sequence as required in the particular embodiment.

The primers shown in Table 2 or 7 may also be used in the invention as mentioned herein. In one embodiment primers are used which comprise any of: the sequences shown in Table 2 or 7; or fragments and/or homologues of any sequence shown in Table 2 or 7.

The Second Set of Nucleic Acids – the 'Index' Sequences

The second set of nucleic acid sequences has the function of being a set of index sequences, and is essentially a set of nucleic acid sequences which are suitable for identifying subgroup specific sequence. They can represent the 'background' chromosomal interactions and might be selected in some way or

35 be unselected. They are in general a subset of all possible chromosomal interactions.

The second set of nucleic acids may be derived by any suitable process. They can be derived computationally or they may be based on chromosome interaction in individuals. They typically represent a larger population group than the first set of nucleic acids. In one particular embodiment, the second set

5 of nucleic acids represents all possible epigenetic chromosomal interactions in a specific set of genes. In another particular embodiment, the second set of nucleic acids represents a large proportion of all possible epigenetic chromosomal interactions present in a population described herein. In one particular embodiment, the second set of nucleic acids represents at least 50% or at least 80% of epigenetic chromosomal interactions in at least 20, 50, 100 or 500 genes, for example in 20 to 100 or 50 to 500 genes.

10 The second set of nucleic acids typically represents at least 100 possible epigenetic chromosome interactions which modify, regulate or in any way mediate a disease state / phenotype in population. The second set of nucleic acids may represent chromosome interactions that affect a disease state (typically relevant to diagnosis or prognosis) in a species. The second set of nucleic acids typically comprises

15 sequences representing epigenetic interactions both relevant and not relevant to an ALS subgroup.

In one particular embodiment the second set of nucleic acids derive at least partially from naturally occurring sequences in a population, and are typically obtained by *in silico* processes. Said nucleic acids may further comprise single or multiple mutations in comparison to a corresponding portion of nucleic acids present in the naturally occurring nucleic acids. Mutations include deletions, substitutions and/or

20 additions of one or more nucleotide base pairs. In one particular embodiment, the second set of nucleic acids may comprise sequence representing a homologue and/or orthologue with at least 70% sequence identity to the corresponding portion of nucleic acids present in the naturally occurring species. In another particular embodiment, at least 80% sequence identity or at least 90% sequence identity to the corresponding portion of nucleic acids present in the naturally occurring species is provided.

25 *Properties of the Second Set of Nucleic Acids*

In one particular embodiment, there are at least 100 different nucleic acid sequences in the second set of nucleic acids, preferably at least 1000, 2000 or 5000 different nucleic acids sequences, with up to 100,000, 1,000,000 or 10,000,000 different nucleic acid sequences. A typical number would be 100 to 1,000,000,

30 such as 1,000 to 100,000 different nucleic acids sequences. All or at least 90% or at least 50% of these would correspond to different chromosomal interactions.

In one particular embodiment, the second set of nucleic acids represent chromosome interactions in at least 20 different loci or genes, preferably at least 40 different loci or genes, and more preferably at least

35 100, at least 500, at least 1000 or at least 5000 different loci or genes, such as 100 to 10,000 different loci or genes. The lengths of the second set of nucleic acids are suitable for them to specifically hybridise

according to Watson Crick base pairing to the first set of nucleic acids to allow identification of chromosome interactions specific to subgroups. Typically the second set of nucleic acids will comprise two portions corresponding in sequence to the two chromosome regions which come together in the chromosome interaction. The second set of nucleic acids typically comprise nucleic acid sequences which

5 are at least 10, preferably 20, and preferably still 30 bases (nucleotides) in length. In another embodiment, the nucleic acid sequences may be at the most 500, preferably at most 100, and preferably still at most 50 base pairs in length. In a preferred embodiment, the second set of nucleic acids comprises nucleic acid sequences of between 17 and 25 base pairs. In one embodiment at least 100, 80% or 50% of the second set of nucleic acid sequences have lengths as described above. Preferably the different nucleic acids do

10 not have any overlapping sequences, for example at least 100%, 90%, 80% or 50% of the nucleic acids do not have the same sequence over at least 5 contiguous nucleotides.

Given that the second set of nucleic acids acts as an 'index' then the same set of second nucleic acids may be used with different sets of first nucleic acids which represent subgroups for different characteristics,

15 i.e. the second set of nucleic acids may represent a 'universal' collection of nucleic acids which can be used to identify chromosome interactions relevant to different characteristics.

The First Set of Nucleic Acids

The first set of nucleic acids are typically from subgroups relevant to diagnosis or prognosis of ALS or

20 Huntington's disease. The first nucleic acids may have any of the characteristics and properties of the second set of nucleic acids mentioned herein. The first set of nucleic acids is normally derived from a sample from the individuals which has undergone treatment and processing as described herein, particularly the EpiSwitch™ cross-linking and cleaving steps. Typically the first set of nucleic acids represents all or at least 80% or 50% of the chromosome interactions present in the samples taken from

25 the individuals.

Typically, the first set of nucleic acids represents a smaller population of chromosome interactions across the loci or genes represented by the second set of nucleic acids in comparison to the chromosome interactions represented by second set of nucleic acids, i.e. the second set of nucleic acids is representing

30 a background or index set of interactions in a defined set of loci or genes.

Library of Nucleic Acids

Any of the types of nucleic acid populations mentioned herein may be present in the form of a library comprising at least 200, at least 500, at least 1000, at least 5000 or at least 10000 different nucleic acids

of that type, such as 'first' or 'second' nucleic acids. Such a library may be in the form of being bound to an array.

Hybridisation

5 The invention requires a means for allowing wholly or partially complementary nucleic acid sequences from the first set of nucleic acids and the second set of nucleic acids to hybridise. In one embodiment all of the first set of nucleic acids is contacted with all of the second set of nucleic acids in a single assay, i.e. in a single hybridisation step. However any suitable assay can be used.

10 *Labelled Nucleic Acids and Pattern of Hybridisation*

The nucleic acids mentioned herein may be labelled, preferably using an independent label such as a fluorophore (fluorescent molecule) or radioactive label which assists detection of successful hybridisation. Certain labels can be detected under UV light. The pattern of hybridisation, for example on an array described herein, represents differences in epigenetic chromosome interactions between the two 15 subgroups, and thus provides a process of comparing epigenetic chromosome interactions and determination of which epigenetic chromosome interactions are specific to a subgroup in the population of the present invention.

The term 'pattern of hybridisation' broadly covers the presence and absence of hybridisation between 20 the first and second set of nucleic acids, i.e. which specific nucleic acids from the first set hybridise to which specific nucleic acids from the second set, and so it not limited to any particular assay or technique, or the need to have a surface or array on which a 'pattern' can be detected.

Selecting a Subgroup with Particular Characteristics

25 The invention provides a process which comprises detecting the presence or absence of chromosome interactions, typically 5 to 20 or 5 to 500 such interactions, preferably 20 to 300 or 50 to 100 interactions, in order to determine the presence or absence of a characteristic relating to ALS or Huntington's disease in an individual. Preferably the chromosome interactions are those in any of the genes mentioned herein. In one embodiment the chromosome interactions which are typed are those represented by the nucleic 30 acids in Table 1 or 5. Preferably the chromosome interactions which are typed are those represented by the nucleic acids in Table 10, 11 or 12. The column titled 'Loop Detected' in the tables shows which subgroup is detected (ALS or control) by each probe. As can be seen the process of the invention can detect either an ALS subgroup and/or a control subgroup (non-ALS) as part of the testing.

The Individual that is Tested

Examples of the species that the individual who is tested is from are mentioned herein. In addition the individual that is tested in the process of the invention may have been selected in some way. The individual may be susceptible to ALS or Huntington's disease, for example.

5

Preferred Gene Regions, Loci, Genes and Chromosome Interactions

For all aspects of the invention preferred gene regions, loci, genes and chromosome interactions are mentioned in the tables, for example in Table 1 and 5. Typically in the processes of the invention chromosome interactions are detected from at least 1, 2, 3, 4, 5, 6, 7 or 8 of the relevant genes listed in

10 Table 1 or 5. Preferably the presence or absence of at least 1, 2, 3, 4, 5, 6, 7 or 8 of the relevant specific chromosome interactions represented by the probe sequences in Table 1 or 5 are detected. The chromosome interaction may be upstream or downstream of any of the genes mentioned herein, for example 50 kb upstream or 20 kb downstream, for example from the coding sequence.

Preferably at least 5, 8, 10, 15 or all of the chromosome interactions in Table 1 are typed.

15 Typically embodiment at least 5, 7, 8 or all of the chromosome interactions in Table 5 are typed.

Preferably at least 4, 6 or all of the chromosome interactions in Table 10 are typed.

Typically at least 1, 2 or all of the chromosome interactions in Table 11 are typed.

Preferably at least 4, 6 or all of the chromosome interactions in Table 12 are typed.

Chromosome interactions may be typed to determine presence of disease. Chromosome interactions may

20 be typed to determine whether the individual will progress fast or slow.

The specific probes and primer sequence disclosed in the tables (or derivates including fragments and homologues) may be used for any typing method disclosed herein. Their use in a method of diagnosis or prognosis is provided.

25 In one embodiment the locus (including the gene and/or place where the chromosome interaction is detected) may comprise a CTCF binding site. This is any sequence capable of binding transcription repressor CTCF. That sequence may consist of or comprise the sequence CCCTC which may be present in 1, 2 or 3 copies at the locus. The CTCF binding site sequence may comprise the sequence CCGCGNGGNGGCAG (in IUPAC notation). The CTCF binding site may be within at least 100, 500, 1000 or 4000 bases of the chromosome interaction or within any of the chromosome regions shown Table 1 or 5.

In one embodiment the chromosome interactions which are detected are present at any of the gene regions shown Table 1 or 5. In the case where a ligated nucleic acid is detected in the process then sequence shown in any of the probe sequences in Table 1 or 5 may be detected.

Thus typically sequence from both regions of the probe (i.e. from both sites of the chromosome interaction) could be detected. In preferred embodiments probes are used in the process which comprise or consist of the same or complementary sequence to a probe shown in any table. In some embodiments probes are used which comprise sequence which is homologous to any of the probe sequences shown in the tables.

Tables Provided Herein

10 Tables 1 and 5 show probe (Episwitch™ marker) data and gene data representing chromosome interactions relevant to ALS. The probe sequences show sequence which can be used to detect a ligated product generated from both sites of gene regions that have come together in chromosome interactions, i.e. the probe will comprise sequence which is complementary to sequence in the ligated product. The first two sets of Start-End positions show probe positions, and the second two sets of Start-End positions
15 show the relevant 4kb region. The following information is provided in the probe data table:

- HyperG_Stats: p-value for the probability of finding that number of significant EpiSwitch™ markers in the locus based on the parameters of hypergeometric enrichment
- Probe Count Total: Total number of EpiSwitch™ Conformations tested at the locus
- Probe Count Sig: Number of EpiSwitch™ Conformations found to be statistical significant at the locus
- FDR HyperG: Multi-test (False Discovery Rate) corrected hypergeometric p-value
- Percent Sig: Percentage of significant EpiSwitch™ markers relative the number of markers tested at the locus
- logFC: logarithm base 2 of Epigenetic Ratio (FC)
- AveExpr: average log2-expression for the probe over all arrays and channels
- T: moderated t-statistic
- p-value: raw p-value
- adj. p-value: adjusted p-value or q-value
- B - B-statistic (lods or B) is the log-odds that that gene is differentially expressed.
- FC - non-log Fold Change
- FC_1 - non-log Fold Change centred around zero
- LS – Binary value this relates to FC_1 values. FC_1 value below -1.1 it is set to -1 and if the FC_1 value is above 1.1 it is set to 1. Between those values the value is 0

Tables 1 and 5 show genes where a relevant chromosome interaction has been found to occur. Other
35 tables show similar data. The p-value in the loci table is the same as the HyperG_Stats (p-value for the probability of finding that number of significant EpiSwitch™ markers in the locus based on the parameters of hypergeometric enrichment).

The probes are designed to be 30bp away from the Taq1 site. In case of PCR, PCR primers are also designed to detect ligated product but their locations from the Taq1 site vary.

Probe locations:

Start 1 - 30 bases upstream of Taq1 site on fragment 1

5 End 1 - Taq1 restriction site on fragment 1

Start 2 - Taq1 restriction site on fragment 2

End 2 - 30 bases downstream of Taq1 site on fragment 2

4kb Sequence Location:

Start 1 - 4000 bases upstream of Taq1 site on fragment 1

10 End 1 - Taq1 restriction site on fragment 1

Start 2 - Taq1 restriction site on fragment 2

End 2 - 4000 bases downstream of Taq1 site on fragment 2

GLMNET values related to procedures for fitting the entire lasso or elastic-net regularization (Lambda set to 0.5 (elastic-net)).

15 Tables 1 to 4 relate to diagnosis of ALS, whilst Tables 5 to 9 relate to prognosis of ALS, and in one embodiment detection relevant to diagnosis is performed based on Tables 1 and 4 and detection relevant to prognosis is performed based on Tables 5 to 9.

Table 1 - The LS column shows a 1 or -1. 1 means present in ALS cases and -1 means absent from ALS cases.

20 Table 5 - The LS column shows 1 or -1. 1 means the marker is present in fast progressors and absent from slow progressors, and -1 means the marker is present in slow progressors but absent in fast progressors.

Tables 10 and 11 relate to ALS prognosis. Table 10 includes markers shown in earlier tables. The 'Loop detected' column of Table 11 means the marker is present in fast progressors and absent from slow progressors.

25 Markers are uniquely identified in the tables with reference to a relevant probe for the product of an EpiSwitch 3C method. In the case of hydrolysis probes these are located on top of Taq site (TCGA) and they cover both genome sites in EpiSwitch interaction . It measures the same junction as 60 base

array probes (30 bases on each side of the sequence tags), but spans over a tailored length of the sequence on both sides. Table 19 provides an example of this.

Preferred Embodiments for Sample Preparation and Chromosome Interaction Detection

5 Methods of preparing samples and detecting chromosome conformations are described herein. Optimised (non-conventional) versions of these methods can be used, for example as described in this section.

Typically the sample will contain at least 2×10^5 cells. The sample may contain up to 5×10^5 cells. In one embodiment, the sample will contain 2×10^5 to 5.5×10^5 cells

10 Crosslinking of epigenetic chromosomal interactions present at the chromosomal locus is described herein. This may be performed before cell lysis takes place. Cell lysis may be performed for 3 to 7 minutes, such as 4 to 6 or about 5 minutes. In some embodiments, cell lysis is performed for at least 5 minutes and for less than 10 minutes.

15 Digesting DNA with a restriction enzyme is described herein. Typically, DNA restriction is performed at about 55°C to about 70°C , such as for about 65°C , for a period of about 10 to 30 minutes, such as about 20 minutes.

Preferably a frequent cutter restriction enzyme is used which results in fragments of ligated DNA with an average fragment size up to 4000 base pair. Optionally the restriction enzyme results in fragments of ligated DNA have an average fragment size of about 200 to 300 base pairs, such as about 256 base pairs.

20 In one embodiment, the typical fragment size is from 200 base pairs to 4,000 base pairs, such as 400 to 2,000 or 500 to 1,000 base pairs.

In one embodiment of the EpiSwitch method a DNA precipitation step is not performed between the DNA restriction digest step and the DNA ligation step.

25 DNA ligation is described herein. Typically the DNA ligation is performed for 5 to 30 minutes, such as about 10 minutes.

The protein in the sample may be digested enzymatically, for example using a proteinase, optionally Proteinase K. The protein may be enzymatically digested for a period of about 30 minutes to 1 hour, for example for about 45 minutes. In one embodiment after digestion of the protein, for example Proteinase K digestion, there is no cross-link reversal or phenol DNA extraction step.

30 In one embodiment PCR detection is capable of detecting a single copy of the ligated nucleic acid, preferably with a binary read-out for presence/absence of the ligated nucleic acid.

Processes and Uses of the Invention

The process of the invention can be described in different ways. It can be described as a method of making a ligated nucleic acid comprising (i) *in vitro* cross-linking of chromosome regions which have come together in a chromosome interaction; (ii) subjecting said cross-linked DNA to cutting or restriction digestion cleavage; and (iii) ligating said cross-linked cleaved DNA ends to form a ligated nucleic acid, wherein detection of the ligated nucleic acid may be used to determine the chromosome state at a locus, and wherein preferably:

- the locus may be any of the loci, regions or genes mentioned in Table 1 or 5,
- 10 - and/or wherein the chromosomal interaction may be any of the chromosome interactions mentioned herein or corresponding to any of the probes disclosed in Table 1 or 5, and/or
- wherein the ligated product may have or comprise (i) sequence which is the same as or homologous to any of the probe sequences disclosed in Table 1 or 5; or (ii) sequence which is complementary to (ii).

The process of the invention can be described as a process for detecting chromosome states which represent different subgroups in a population comprising determining whether a chromosome interaction is present or absent within a defined epigenetically active (disease associated) region of the genome, wherein preferably:

- the subgroup is defined by presence or absence of ALS or a characteristic relating to ALS (such as prognosis or progression), and/or
- 20 - the chromosome state may be at any locus, region or gene mentioned in Table 1 or 5; and/or
- the chromosome interaction may be any of those mentioned in Table 1 or 5 or corresponding to any of the probes disclosed in that table.

The invention includes detecting chromosome interactions at any locus, gene or regions mentioned Table 1 or 5. The invention includes use of the nucleic acids and probes mentioned herein to detect chromosome interactions, for example use of at least 1, 2, 4, 6 or 8 such nucleic acids or probes to detect chromosome interactions in at least 1, 2, 4, 6 or 8 different loci or genes. The invention includes detection of chromosome interactions using any of the primers or primer pairs listed in Table 2 or 7 or using variants of these primers as described herein (sequences comprising the primer sequences or comprising fragments and/or homologues of the primer sequences).

When analysing whether a chromosome interaction occurs 'within' a defined gene, region or location, either both the parts of the chromosome which have together in the interaction are within the defined

gene, region or location or in some embodiments only one part of the chromosome is within the defined, gene, region or location.

Use of the Method of the Invention to Identify New Treatments

5 Knowledge of chromosome interactions can be used to identify new treatments for conditions. The invention provides methods and uses of chromosomes interactions defined here to identify or design new therapeutic agents for ALS (including treatments relating to prognosis).

Homologues

Homologues of polynucleotide / nucleic acid (e.g. DNA) sequences are referred to herein. Such

10 homologues typically have at least 70% homology, preferably at least 80%, at least 85%, at least 90%, at least 95%, at least 97%, at least 98% or at least 99% homology, for example over a region of at least 10, 15, 20, 30, 100 or more contiguous nucleotides, or across the portion of the nucleic acid which is from the region of the chromosome involved in the chromosome interaction. The homology may be calculated on the basis of nucleotide identity (sometimes referred to as "hard homology").

15 Therefore, in a particular embodiment, homologues of polynucleotide / nucleic acid (e.g. DNA) sequences are referred to herein by reference to percentage sequence identity. Typically such homologues have at least 70% sequence identity, preferably at least 80%, at least 85%, at least 90%, at least 95%, at least 97%, at least 98% or at least 99% sequence identity, for example over a region of at least 10, 15, 20, 30, 100 or more contiguous nucleotides, or across the portion of the nucleic acid which

20 is from the region of the chromosome involved in the chromosome interaction.

For example the UWGCG Package provides the BESTFIT program which can be used to calculate homology and/or % sequence identity (for example used on its default settings) (Devereux *et al* (1984) Nucleic Acids Research 12, p387-395). The PILEUP and BLAST algorithms can be used to calculate homology and/or % sequence identity and/or line up sequences (such as identifying equivalent or

25 corresponding sequences (typically on their default settings)), for example as described in Altschul S. F. (1993) J Mol Evol 36:290-300; Altschul, S, F *et al* (1990) J Mol Biol 215:403-10.

Software for performing BLAST analyses is publicly available through the National Center for Biotechnology Information. This algorithm involves first identifying high scoring sequence pair (HSPs) by identifying short words of length W in the query sequence that either match or satisfy some positive-valued threshold score T when aligned with a word of the same length in a database sequence. T is referred to as the neighbourhood word score threshold (Altschul *et al*, *supra*). These initial neighbourhood word hits act as seeds for initiating searches to find HSPs containing them. The word hits are extended in both directions along each sequence for as far as the cumulative alignment score can be increased. Extensions for the word hits in each direction are halted when: the cumulative alignment score falls off

by the quantity X from its maximum achieved value; the cumulative score goes to zero or below, due to the accumulation of one or more negative-scoring residue alignments; or the end of either sequence is reached. The BLAST algorithm parameters W5 T and X determine the sensitivity and speed of the alignment. The BLAST program uses as defaults a word length (W) of 11, the BLOSUM62 scoring matrix

5 (see Henikoff and Henikoff (1992) Proc. Natl. Acad. Sci. USA 89: 10915-10919) alignments (B) of 50, expectation (E) of 10, M=5, N=4, and a comparison of both strands.

The BLAST algorithm performs a statistical analysis of the similarity between two sequences; see e.g., Karlin and Altschul (1993) Proc. Natl. Acad. Sci. USA 90: 5873-5787. One measure of similarity provided by the BLAST algorithm is the smallest sum probability (P(N)), which provides an indication of the

10 probability by which a match between two polynucleotide sequences would occur by chance. For example, a sequence is considered similar to another sequence if the smallest sum probability in comparison of the first sequence to the second sequence is less than about 1, preferably less than about 0.1, more preferably less than about 0.01, and most preferably less than about 0.001.

15 The homologous sequence typically differs by 1, 2, 3, 4 or more bases, such as less than 10, 15 or 20 bases (which may be substitutions, deletions or insertions of nucleotides). These changes may be measured across any of the regions mentioned above in relation to calculating homology and/or % sequence identity.

20 Homology of a 'pair of primers' can be calculated, for example, by considering the two sequences as a single sequence (as if the two sequences are joined together) for the purpose of then comparing against the another primer pair which again is considered as a single sequence.

Arrays

The second set of nucleic acids may be bound to an array, and in one embodiment there are at least 15,000, 45,000, 100,000 or 250,000 different second nucleic acids bound to the array, which preferably 25 represent at least 300, 900, 2000 or 5000 loci. In one embodiment one, or more, or all of the different populations of second nucleic acids are bound to more than one distinct region of the array, in effect repeated on the array allowing for error detection. The array may be based on an Agilent SurePrint G3 Custom CGH microarray platform. Detection of binding of first nucleic acids to the array may be performed by a dual colour system.

30

Therapeutic Agents

Therapeutic agents are mentioned herein. The invention provides such agents for use in preventing or treating ALS in certain individuals, for example those identified by a process of the invention. This may comprise administering to an individual in need a therapeutically effective amount of the agent. The

invention provides use of the agent in the manufacture of a medicament to prevent or treat ALS in certain individuals.

Preferred therapeutic agents for ALS are:

Riluzole (Rilutek): This drug is typically taken as a pill and it reduces the disease's progression by

5 reducing levels of a messenger (glutamate) in the brain. Glutamate is present in higher levels in ALS patient.

Edaravone (Radicava): This drug reduces the decline in daily performance related to ALS. The drug is typically given to patient via intravenous infusion for 10-14 days in a row, once a month.

Arimoclomol: This drug performs as a heat shock response inducer in motor neurons and it defends 10 against neuronal disorder and cell death.

Talampanel: This drug reduces the rate of muscle strength decline and symptoms progression.

Beta-lactam antibiotics: These antibiotics such as penicillin and cephalosporin maintain muscle stability and increase life time via up-regulating the level of GLT1, glial glutamate transporter 1.

Bromocriptine: This drug is a free-radical scavenger that inhibits oxidative cell death induced by stress.

15 **Pramipexole and Dexpramipexole:** These drugs act as a dopamine agonist and they have a free-radical scavenging function. These drugs are involved in mitochondrial malfunction. Dexpramipexole is an optical enantiomer of pramipexole.

Stem Cell therapy: Stem cell growth reduces the progression of neuron disease or replaces motor neurons. Stem cells have the potential to make spinal motor neurons, expand axons and receive and 20 generate synapses with the muscle. Mesenchymal stem cells (MSCs) derived from adult stem cells releases trophic factors, anti-inflammatory cytokines and immunomodulatory chemokines to postpone disease progression.

Immunotherapy: Antibody therapy, such as D3H5 antibody infusion via ICV route maintains weight for a longer period of time and prolongs the life time of transgenic mouse model for ALS.

25 The following is a list of therapies for Huntington's disease. These can help reduce some symptoms of movement and psychiatric disorders.

Medications for movement disorders

- Tetrabenazine (Xenazine)
- Antipsychotic drugs, such as haloperidol (Haldol), chlorpromazine, risperidone (Risperdal) and 30 quetiapine (Seroquel)

- Other medications include amantadine, levetiracetam (Keppra, others) and clonazepam (Klonopin)

Medications for psychiatric disorders

- Antidepressants include citalopram (Celexa), escitalopram (Lexapro), fluoxetine (Prozac, Sarafem) and sertraline (Zoloft)
- Antipsychotics include quetiapine (Seroquel), risperidone (Risperdal) and olanzapine (Zyprexa)
- Mood-stabilizing drugs include valproate (Depacon), carbamazepine (Carbatrol, Epitol, Tegretol) and lamotrigine (Lamictal)

10 Early diagnosis of Huntington's helps to manage the treatment of symptoms over the course of the disease.

The formulation of the agent (for ALS or Huntington's disease) will depend upon the nature of the agent. The agent will be provided in the form of a pharmaceutical composition containing the agent and a pharmaceutically acceptable carrier or diluent. Suitable carriers and diluents include isotonic saline 15 solutions, for example phosphate-buffered saline. Typical oral dosage compositions include tablets, capsules, liquid solutions and liquid suspensions. The agent may be formulated for parenteral, intravenous, intramuscular, subcutaneous, transdermal or oral administration.

20 The dose of an agent may be determined according to various parameters, especially according to the substance used; the age, weight and condition of the individual to be treated; the route of administration; and the required regimen. A physician will be able to determine the required route of administration and dosage for any particular agent. A suitable dose may however be from 0.1 to 100 mg/kg body weight such as 1 to 40 mg/kg body weight, for example, to be taken from 1 to 3 times daily.

Forms of the Substance Mentioned Herein

25 Any of the substances, such as nucleic acids or therapeutic agents, mentioned herein may be in purified or isolated form. They may be in a form which is different from that found in nature, for example they may be present in combination with other substance with which they do not occur in nature. The nucleic acids (including portions of sequences defined herein) may have sequences which are different to those found in nature, for example having at least 1, 2, 3, 4 or more nucleotide changes in the sequence as 30 described in the section on homology. The nucleic acids may have heterologous sequence at the 5' or 3' end. The nucleic acids may be chemically different from those found in nature, for example they may be modified in some way, but preferably are still capable of Watson-Crick base pairing. Where appropriate

the nucleic acids will be provided in double stranded or single stranded form. The invention provides all of the specific nucleic acid sequences mentioned herein in single or double stranded form, and thus includes the complementary strand to any sequence which is disclosed.

The invention also provides a kit for carrying out any process of the invention, including detection of a chromosomal interaction associated with ALS (relating for example to diagnosis or prognosis of ALS). Such a kit can include a specific binding agent capable of detecting the relevant chromosomal interaction, such as agents capable of detecting a ligated nucleic acid generated by processes of the invention. Preferred agents present in the kit include probes capable of hybridising to the ligated nucleic acid or primer pairs, for example as described herein, capable of amplifying the ligated nucleic acid in a PCR reaction.

10 The invention also provides a device that is capable of detecting the relevant chromosome interactions. The device preferably comprises any specific binding agents, probe or primer pair capable of detecting the chromosome interaction, such as any such agent, probe or primer pair described herein.

Detection Methods

15 In one embodiment quantitative detection of the ligated sequence which is relevant to a chromosome interaction is carried out using a probe which is detectable upon activation during a PCR reaction, wherein said ligated sequence comprises sequences from two chromosome regions that come together in an epigenetic chromosome interaction, wherein said method comprises contacting the ligated sequence with the probe during a PCR reaction, and detecting the extent of activation of the probe, and wherein said probe binds the ligation site. The method typically allows particular interactions to be

20 detected in a MIQE compliant manner using a dual labelled fluorescent hydrolysis probe.

The probe is generally labelled with a detectable label which has an inactive and active state, so that it is only detected when activated. The extent of activation will be related to the extent of template (ligation product) present in the PCR reaction. Detection may be carried out during all or some of the PCR, for example for at least 50% or 80% of the cycles of the PCR.

25 The probe can comprise a fluorophore covalently attached to one end of the oligonucleotide, and a quencher attached to the other end of the nucleotide, so that the fluorescence of the fluorophore is quenched by the quencher. In one embodiment the fluorophore is attached to the 5' end of the oligonucleotide, and the quencher is covalently attached to the 3' end of the oligonucleotide. Fluorophores that can be used in the methods of the invention include FAM, TET, JOE, Yakima Yellow,

30 HEX, Cyanine3, ATTO 550, TAMRA, ROX, Texas Red, Cyanine 3.5, LC610, LC 640, ATTO 647N, Cyanine 5, Cyanine 5.5 and ATTO 680. Quenchers that can be used with the appropriate fluorophore include TAM, BHQ1, DAB, Eclip, BHQ2 and BBQ650, optionally wherein said fluorophore is selected from HEX, Texas

Red and FAM. Preferred combinations of fluorophore and quencher include FAM with BHQ1 and Texas Red with BHQ2.

Use of the Probe in a qPCR Assay

Hydrolysis probes of the invention are typically temperature gradient optimised with concentration matched negative controls. Preferably single-step PCR reactions are optimized. More preferably a standard curve is calculated. An advantage of using a specific probe that binds across the junction of the ligated sequence is that specificity for the ligated sequence can be achieved without using a nested PCR approach. The methods described herein allow accurate and precise quantification of low copy number targets. The target ligated sequence can be purified, for example gel-purified, prior to temperature gradient optimization. The target ligated sequence can be sequenced. Preferably PCR reactions are performed using about 10ng, or 5 to 15 ng, or 10 to 20ng, or 10 to 50ng, or 10 to 200ng template DNA. Forward and reverse primers are designed such that one primer binds to the sequence of one of the chromosome regions represented in the ligated DNA sequence, and the other primer binds to other chromosome region represented in the ligated DNA sequence, for example, by being complementary to the sequence.

Choice of Ligated DNA Target

The invention includes selecting primers and a probe for use in a PCR method as defined herein comprising selecting primers based on their ability to bind and amplify the ligated sequence and selecting the probe sequence based properties of the target sequence to which it will bind, in particular the curvature of the target sequence.

Probes are typically designed/chosen to bind to ligated sequences which are juxtaposed restriction fragments spanning the restriction site. In one embodiment of the invention, the predicted curvature of possible ligated sequences relevant to a particular chromosome interaction is calculated, for example using a specific algorithm referenced herein. The curvature can be expressed as degrees per helical turn, e.g. 10.5° per helical turn. Ligated sequences are selected for targeting where the ligated sequence has a curvature propensity peak score of at least 5° per helical turn, typically at least 10°, 15° or 20° per helical turn, for example 5° to 20° per helical turn. Preferably the curvature propensity score per helical turn is calculated for at least 20, 50, 100, 200 or 400 bases, such as for 20 to 400 bases upstream and/or downstream of the ligation site. Thus in one embodiment the target sequence in the ligated product has any of these levels of curvature. Target sequences can also be chosen based on lowest thermodynamic structure free energy.

Preferred ALS Embodiments

Paragraph 1. A process for detecting a chromosome state which represents a disease subgroup in a population comprising determining whether a chromosome interaction relating to that chromosome

state is present or absent within a defined region of the genome, wherein said disease subgroup is an amyotrophic lateral sclerosis (ALS) subgroup; and

- wherein said chromosome interaction has optionally been identified by a method of determining which chromosomal interactions are relevant to a chromosome state corresponding to an ALS subgroup

5 of the population, comprising contacting a first set of nucleic acids from subgroups with different states of the chromosome with a second set of index nucleic acids, and allowing complementary sequences to hybridise, wherein the nucleic acids in the first and second sets of nucleic acids represent a ligated product comprising sequences from both the chromosome regions that have come together in chromosomal interactions, and wherein the pattern of hybridisation between the first and second set of 10 nucleic acids allows a determination of which chromosomal interactions are specific to an ALS subgroup; and

- wherein the chromosome interaction:

(i) is present in any one of the regions or genes listed in Table 1 or 5; and/or

15 (ii) corresponds to any one of the chromosome interactions represent by any probe shown in Table 1 or 5, and/or

(iii) is present in a 4,000 base region which comprises or which flanks (i) or (ii).

Paragraph 2. A process according to paragraph 1 which is carried out to diagnose ALS or to determine a prognosis for ALS.

20 Paragraph 3. A process according to paragraph 1 or 2 wherein a specific combination of chromosome interactions are typed:

(i) comprising all of the chromosome interactions represented by the probes in Table 1 or 5; or

(ii) comprising at least 4, 5, 6 or 7 of the chromosome interactions represented by the probes in Table 1 or 5; or

(iii) which together are present in at least 4, 5, 6 or 7 of the regions or genes listed in Table 1 or 5; or

25 (iv) at least 4, 5, 6 or 7 chromosome interactions are typed which are present in a 4,000 base region which comprises or which flanks the chromosome interactions represented by the probes in Table 1 or 5.

Paragraph 4. A process according to any one of the preceding paragraphs in which the chromosome interactions are typed:

30 - in a sample from an individual, and/or

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- by detecting the presence or absence of a DNA loop at the site of the chromosome interactions, and/or
- detecting the presence or absence of distal regions of a chromosome being brought together in a chromosome conformation, and/or

5 - by detecting the presence of a ligated nucleic acid which is generated during said typing and whose sequence comprises two regions each corresponding to the regions of the chromosome which come together in the chromosome interaction, wherein detection of the ligated nucleic acid is preferably by using (i) a probe that has at least 70% identity to any of the specific probe sequences mentioned in Table 1 or 5, and/or (ii) by a primer pair which has at least 70% identity to any primer pair in Table 2 or 7.

10 Paragraph 5. A process according to any one of the preceding paragraphs, wherein:

- the second set of nucleic acids is from a larger group of individuals than the first set of nucleic acids; and/or
- the first set of nucleic acids is from at least 8 individuals; and/or
- the first set of nucleic acids is from at least 4 individuals from a first subgroup and at least 4 individuals from a second subgroup which is preferably non-overlapping with the first subgroup; and/or
- the process is carried out to select an individual for a medical treatment.

Paragraph 6. A process according to any one of the preceding paragraphs wherein:

15 - the second set of nucleic acids represents an unselected group; and/or

- wherein the second set of nucleic acids is bound to an array at defined locations; and/or

20 - wherein the second set of nucleic acids represents chromosome interactions in at least 100 different genes; and/or

- wherein the second set of nucleic acids comprises at least 1,000 different nucleic acids representing at least 1,000 different chromosome interactions; and/or

25 - wherein the first set of nucleic acids and the second set of nucleic acids comprise at least 100 nucleic acids with length 10 to 100 nucleotide bases.

Paragraph 7. A process according to any one of the preceding paragraphs, wherein the first set of nucleic acids is obtainable in a process comprising the steps of: -

- (i) cross-linking of chromosome regions which have come together in a chromosome interaction;

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- (ii) subjecting said cross-linked regions to cleavage, optionally by restriction digestion cleavage with an enzyme; and
- (iii) ligating said cross-linked cleaved DNA ends to form the first set of nucleic acids (in particular comprising ligated DNA).

5 Paragraph 8. A process according to any one of the preceding paragraphs wherein at least 5 to 9 different chromosome interactions are typed, preferably in 5 to 9 different regions or genes.

Paragraph 9. A process according to any one of the preceding paragraphs wherein said defined region of the genome:

- (i) comprises a single nucleotide polymorphism (SNP); and/or
- 10 (ii) expresses a microRNA (miRNA); and/or
- (iii) expresses a non-coding RNA (ncRNA); and/or
- (iv) expresses a nucleic acid sequence encoding at least 10 contiguous amino acid residues; and/or
- (v) expresses a regulating element; and/or
- (vii) comprises a CTCF binding site.

15 Paragraph 10. Method for identifying or designing a therapeutic agent for treating ALS by selecting an agent that is able to cause a change of chromosomal interaction and thereby cause a therapeutic effect,

- wherein the chromosomal interaction is represented by any probe in Table 1 or 5; and/or
- the chromosomal interaction is present in any region or gene listed in Table 1 or 5;

and wherein optionally:

- 20 - the chromosomal interaction has been identified by the method of determining which chromosomal interactions are relevant to a chromosome state as defined in paragraph 1, and/or
- the change in chromosomal interaction is monitored using (i) a probe that has at least 70% identity to any of the probe sequences mentioned in Table 1 or 5, and/or (ii) by a primer pair which has at least 70% identity to any primer pair in Table 1 or 5; and/or
- 25 - a candidate agent is contacted with a cell and the chromosome interaction in the cell is monitored to determine whether the candidate agent is able to treat ALS.

Paragraph 11. Use of

- (i) detection of a chromosomal interaction wherein:

- the chromosomal interaction is as represented by a probe in Table 1 or 5, and/or
- the chromosomal interaction is present in any region or gene which is mentioned in Table 1 or 5; or
- (ii) a probe that has at least 70% identity to any of the probe sequences mentioned in Table 1 or 5, or
- 5 (iii) a primer pair which has at least 70% identity to any primer pair identified in Table 2 or 7; to identify or design a therapeutic agent for ALS.

Paragraph 12. Use according to paragraph 11 to identify a therapeutic agent comprising administering a candidate agent, and using said detection of a chromosomal interaction, said probe or said primer pair to detect whether there is a change in chromosome state to thereby determine whether the candidate 10 agent is a therapeutic agent, wherein the use is optionally performed *in vitro*, preferably in a cell.

Paragraph 13. A therapeutic agent for ALS for use in a method of preventing or treating ALS in an individual that has been identified as being in need of the therapeutic agent by a process according to any one of paragraphs 1 to 9.

15 Paragraph 14. A process, method or use according to any of the preceding paragraphs wherein the typing or detecting comprises specific detection of the ligated product by quantitative PCR (qPCR) which uses primers capable of amplifying the ligated product and a probe which binds the ligation site during the PCR reaction, wherein said probe comprises sequence which is complementary to sequence from each of the chromosome regions that have come together in the chromosome interaction, wherein 20 preferably said probe comprises:

- an oligonucleotide which specifically binds to said ligated product, and/or
- a fluorophore covalently attached to the 5' end of the oligonucleotide, and/or
- a quencher covalently attached to the 3' end of the oligonucleotide, and
- optionally

25 said fluorophore is selected from HEX, Texas Red and FAM; and/or

said probe comprises a nucleic acid sequence of length 10 to 40 nucleotide bases, preferably a length of 20 to 30 nucleotide bases.

Particular Embodiments

In one embodiment only intrachromosomal interactions are typed/detected, and no extrachromosomal interactions (between different chromosomes) are typed/detected.

Publications

The contents of all publications mentioned herein are incorporated by reference into the present specification and may be used to further define the features relevant to the invention.

Specific Embodiments

The EpiSwitch™ platform technology detects epigenetic regulatory signatures of regulatory changes between normal and abnormal conditions at loci. The EpiSwitch™ platform identifies and monitors the fundamental epigenetic level of gene regulation associated with regulatory high order structures of human chromosomes also known as chromosome conformation signatures. Chromosome signatures are a distinct primary step in a cascade of gene deregulation. They are high order biomarkers with a unique set of advantages against biomarker platforms that utilize late epigenetic and gene expression biomarkers, such as DNA methylation and RNA profiling.

15 *EpiSwitch™ Array Assay*

The custom EpiSwitch™ array-screening platforms come in 4 densities of, 15K, 45K, 100K, and 250K unique chromosome conformations, each chimeric fragment is repeated on the arrays 4 times, making the effective densities 60K, 180K, 400K and 1 Million respectively.

20 *Custom Designed EpiSwitch™ Arrays*

The 15K EpiSwitch™ array can screen the whole genome including around 300 loci interrogated with the EpiSwitch™ Biomarker discovery technology. The EpiSwitch™ array is built on the Agilent SurePrint G3 Custom CGH microarray platform; this technology offers 4 densities, 60K, 180K, 400K and 1 Million probes. The density per array is reduced to 15K, 45K, 100K and 250K as each EpiSwitch™ probe is presented as a 25 quadruplicate, thus allowing for statistical evaluation of the reproducibility. The average number of potential EpiSwitch™ markers interrogated per genetic loci is 50; as such the numbers of loci that can be investigated are 300, 900, 2000, and 5000.

EpiSwitch™ Custom Array Pipeline

30 The EpiSwitch™ array is a dual colour system with one set of samples, after EpiSwitch™ library generation, labelled in Cy5 and the other of sample (controls) to be compared/ analyzed labelled in Cy3. The arrays are scanned using the Agilent SureScan Scanner and the resultant features extracted using the Agilent Feature Extraction software. The data is then processed using the EpiSwitch™ array processing scripts in R. The arrays are processed using standard dual colour packages in Bioconductor in R: Limma *. The

normalisation of the arrays is done using the `normalisedWithinArrays` function in Limma * and this is done to the on chip Agilent positive controls and EpiSwitch™ positive controls. The data is filtered based on the Agilent Flag calls, the Agilent control probes are removed and the technical replicate probes are averaged, in order for them to be analysed using Limma *. The probes are modelled based on their difference

5 between the 2 scenarios being compared and then corrected by using False Discovery Rate. Probes with Coefficient of Variation (CV) $\leq 30\%$ that are ≤ -1.1 or ≥ 1.1 and pass the $p \leq 0.1$ FDR p-value are used for further screening. To reduce the probe set further Multiple Factor Analysis is performed using the FactorMineR package in R.

10 * Note: LIMMA is Linear Models and Empirical Bayes Processes for Assessing Differential Expression in Microarray Experiments. Limma is an R package for the analysis of gene expression data arising from microarray or RNA-Seq.

15 The pool of probes is initially selected based on adjusted p-value, FC and CV $< 30\%$ (arbitrary cut off point) parameters for final picking. Further analyses and the final list are drawn based only on the first two parameters (adj. p-value; FC).

The invention is illustrated by the following non-limiting examples.

20 **Example 1**

Statistical Pipeline

EpiSwitch™ screening arrays are processed using the EpiSwitch™ Analytical Package in R in order to select 25 high value EpiSwitch™ markers for translation on to the EpiSwitch™ PCR platform.

Step 1

30 Probes are selected based on their corrected p-value (False Discovery Rate, FDR), which is the product of a modified linear regression model. Probes below p-value ≤ 0.1 are selected and then further reduced by their Epigenetic ratio (ER), probes ER have to be ≤ -1.1 or ≥ 1.1 in order to be selected for further analysis. The last filter is a coefficient of variation (CV), probes have to be below ≤ 0.3 .

Step 2

The top 40 markers from the statistical lists are selected based on their ER for selection as markers for PCR translation. The top 20 markers with the highest negative ER load and the top 20 markers with the highest positive ER load form the list.

Step 3

5 The resultant markers from step 1, the statistically significant probes form the bases of enrichment analysis using hypergeometric enrichment (HE). This analysis enables marker reduction from the significant probe list, and along with the markers from step 2 forms the list of probes translated on to the EpiSwitch™ PCR platform.

10 The statistical probes are processed by HE to determine which genetic locations have an enrichment of statistically significant probes, indicating which genetic locations are hubs of epigenetic difference.

The most significant enriched loci based on a corrected p-value are selected for probe list generation. Genetic locations below p-value of 0.3 or 0.2 are selected. The statistical probes mapping to these genetic locations, with the markers from step 2, form the high value markers for EpiSwitch™ PCR translation.

Array design and processing

15 *Array Design*

1. Genetic loci are processed using the SII software (currently v3.2) to:
 - a. Pull out the sequence of the genome at these specific genetic loci (gene sequence with 50kb upstream and 20kb downstream)
 - b. Define the probability that a sequence within this region is involved in CCs
 - c. Cut the sequence using a specific RE
 - d. Determine which restriction fragments are likely to interact in a certain orientation
 - e. Rank the likelihood of different CCs interacting together.
2. Determine array size and therefore number of probe positions available (x)
3. Pull out x/4 interactions.
- 25 4. For each interaction define sequence of 30bp to restriction site from part 1 and 30bp to restriction site of part 2. Check those regions aren't repeats, if so exclude and take next interaction down on the list. Join both 30bp to define probe.
5. Create list of x/4 probes plus defined control probes and replicate 4 times to create list to be created on array
- 30 6. Upload list of probes onto Agilent Sure design website for custom CGH array.
7. Use probe group to design Agilent custom CGH array.

Array Processing

1. Process samples using EpiSwitch™ Standard Operating Procedure (SOP) for template production.
2. Clean up with ethanol precipitation by array processing laboratory.
3. Process samples as per Agilent SureTag complete DNA labelling kit - Agilent Oligonucleotide Array-based CGH for Genomic DNA Analysis Enzymatic labelling for Blood, Cells or Tissues
- 5 4. Scan using Agilent C Scanner using Agilent feature extraction software.

EpiSwitch™ technology overview

The *EpiSwitch™* platform offers a highly effective means of screening, early detection, companion-diagnosis, monitoring and prognostic analysis of major diseases associated with aberrant and responsive gene expression. The major advantages of this approach is that it is non-invasive, rapid, and relies on 10 highly stable DNA based targets as part of chromosomal signatures, rather than unstable protein/RNA molecules.

EpiSwitch™ biomarker signatures demonstrate high robustness, sensitivity and specificity in the stratification of complex disease phenotypes. This technology takes advantage of the latest breakthroughs 15 in the science of epigenetics, monitoring and evaluation of chromosome conformation signatures as a highly informative class of epigenetic biomarkers. Current research methodologies deployed in academic environment require from 3 to 7 days for biochemical processing of cellular material in order to detect CCSs. Those procedures have limited sensitivity, and reproducibility; and furthermore, do not have the benefit of the targeted insight provided by the *EpiSwitch™ Analytical Package* at the design stage.

20

EpiSwitch™ Array *in silico* marker identification

CCS sites across the genome are directly evaluated by the *EpiSwitch™ Array* on clinical samples from testing cohorts for identification of all relevant stratifying lead biomarkers. The *EpiSwitch™ Array* platform is used for marker identification due to its high-throughput capacity, and its ability to screen 25 large numbers of loci rapidly. The array used was the Agilent custom-CGH array, which allows markers identified through the *in silico* software to be interrogated.

EpiSwitch™ PCR

Potential markers identified by *EpiSwitch™ Array* are then validated either by *EpiSwitch™ PCR* or DNA 30 sequencers (i.e. Roche 454, Nanopore MinION, etc.). The top PCR markers which are statistically significant and display the best reproducibility are selected for further reduction into the final *EpiSwitch™* Signature Set, and validated on an independent cohort of samples. *EpiSwitch™ PCR* can be performed by a trained technician following a standardised operating procedure protocol established. All protocols and manufacture of reagents are performed under ISO 13485 and 9001 accreditation to ensure the quality of 35 the work and the ability to transfer the protocols. *EpiSwitch™ PCR* and *EpiSwitch™ Array* biomarker

platforms are compatible with analysis of both whole blood and cell lines. The tests are sensitive enough to detect abnormalities in very low copy numbers using small volumes of blood.

Analysis of an ALS cohort

5 The inventors have used epigenetic chromosomal interactions as the basis for identifying biomarkers to be used as a companion diagnostic method in ALS. The *EpiSwitch™* biomarker discovery platform was developed by the inventors to detect epigenetic regulatory signature changes such as those driving phenotypic changes implicated in ALS. The *EpiSwitch™* biomarker discovery platform identifies CCSs which define the initial regulatory process in integrating environmental cues into the epigenetic and

10 transcriptional machinery. As such, CCSs are the primary step in a cascade of gene regulation. The CCSs isolated by the *EpiSwitch™* biomarker discovery platform have several well documented advantages: severe biochemical and physiological stability; their binary nature and readout; and their primary position in the eukaryotic cascade of gene regulation.

The ability to detect perturbations in the systemic epigenetic control of gene expression allows both

15 early diagnosis of ALS and establishment of patient prognosis. A comparative interrogation of the cellular regulatory genome architecture from healthy and diseased-patient blood samples revealed two ALS-related epigenetic signatures: one with diagnostic potential and a second for prognosis prediction. For this prospective study, samples collected by the clinical-research group at the Oxford Motor Neuron Disorders Clinic at the Nuffield Department of Clinical Neurosciences (NDCN) at the University of Oxford

20 were analyzed using *EpiSwitch™*, the high throughput platform developed by Oxford BioDynamics for monitoring chromosome conformation signatures. This study compares the clinical annotations of the ALS-FRS-R, Forced Vital Capacity (FVC) and other clinical observations to assign ALS-progression subtypes, with analysis for the epigenetic signatures.

A total of 100 patients, presenting to the Oxford Motor Neuron Disorders Clinic, enrolled in the study

25 and were asked to return at 3 and 6 months. Controls (n=100) were collected. During each visit, participants underwent the ALS-FRS-R and FVC tests, and provided a blood sample. The samples were analyzed to identify either an ALS- disease-related diagnostic signature, or the prognostic disease-related signature (at 3 and 6 months). Results of the clinical assessments were compared to the *EpiSwitch™* analysis at 0, 3 and 6 months. A cut off of a 0.5-point decline per month of the ALS-FRS-R

30 score was used to cluster the ALS patients into progression-subtypes.

Based on the rate of decline in the ALS-FRS-R score, preliminary results from 3 month samples and 6-month samples demonstrated the epigenetic signature selected faster (>0.5) and slower (<0.5) progressing ALS patients with a sensitivity and specificity of 80%. To date, the results indicate the prognostic signature is robust for selecting subtypes of ALS across time. The results shown in the Tables

relate to the diagnostic and prognostic analysis of 75 ALS and 75 control samples (n=150), with an independent sample cohort (n = 50).

Further results (including Table 11) are based on 100 patients with ALS who have been observed and evaluated by standard practice of ALSFRS-R score and FVC (forced vital capacity) at baseline, 3 months 5 and 6 months survival. Based on measurements at baseline and 3 months these patients were classified as slow or fast ALS (standard classification). With our three markers readout at the baseline only, we also classified all the patients as slow or fast (EpiSwitch classification). These patients were then evaluated on the basis of survival at 6 months, and survival for slow and fast groups was compared. Standard classification had significant number of fatal outcomes in the slow group and statistically the 10 survival difference by p value between slow and fast was insignificant, p=0.052. By EpiSwitch classification, the separation between slow and fast at 6 months was highly significant, p=0.0097, and most of the deaths took place in the fast group.

Example 2

Genomic architecture differences at the HTT locus underlie symptomatic and pre-symptomatic cases of 15 Huntington's Disease.

Huntington's disease (HD), a progressive neurodegenerative condition, is a genetic disorder that causes extensive degeneration of neurons in the adult brain, ultimately leading to death. The root cause of HD is an expanded trinucleotide cytosine-adenine-guanine (CAG) repeat in the "huntingtin gene" (HTT), 20 which results in a mutant huntingtin protein with an increased tendency to form intracellular aggregates and cause neuronal death. Importantly, the number of CAG repeats correlate with disease onset and severity and patients with more than 39 CAG repeats will develop HD at some point in their life, the onset of disease can vary by decades within individuals and little is known about this presymptomatic phase. With the advancement of epigenetic approaches and emergence of new technologies for 25 detection of epigenetic regulatory markers for DNA methylation, histone modifications, non-coding RNA and chromosome conformation signatures as part of understanding genome architecture, we were interested in analyzing the systemic regulatory changes in genomic architecture around the HTT gene and its relationship to disease manifestation. Here, we examined the detectable, systemic, non-invasive epigenetic differences at the HTT locus between presymptomatic and symptomatic HD patients 30 compared to unaffected controls.

Methods: Using blood samples from patients with HD and healthy controls we used *EpiSwitch*™, a validated high-resolution industrial platform for the detection of chromosome conformations, to assess chromatin architecture in the immediate vicinity of the HTT gene. We evaluated the absence or 35 presence of conditional, stable chromatin conformations at 20 interaction sites across over 225 kb of

the HTT locus and compared the resulting chromosome conformations between groups of healthy controls, verified symptomatic HD patients (CAG, n>39) and patients with genetically verified CAG extension who had not manifested clinical symptoms of HD.

5 Results: Consistent and stable chromosome conformations were observed across the patient groups when tested in peripheral blood samples. We found two constitutive interactions (occurring in all patient groups and controls) and seven conditional interactions which were present in HD, but not in healthy controls. Most important, we observed three conditional interactions that were present only in HD patients manifesting clinical symptoms (symptomatic cases), but not in presymptomatic cases. 85%
 10 (6 out of 7) of the patients in the symptomatic HD cohort demonstrated at least one of the specific chromosome conformations associated with symptomatic HD.

Conclusion: Our results are the first evidence that the regulatory chromatin architecture at HTT locus is systemically altered in patients with HD. Moreover, the chromatin architecture at the HTT locus also shows conditional differences between clinical stages in symptomatic and presymptomatic HD patients.

15 Given the high clinical utility in having a molecular tool to assess disease progression in HD, these results strongly suggest the non-invasive assessment of systemic chromosome conformation signatures (CCS) can be a valuable addition to prognostic assessment of HD patients.

Introduction

20 Huntington's Disease (HD) is a neurodegenerative condition characterized cellularly by the loss of neurons in the basal ganglia and clinically by uncontrolled movements, emotional problems, and loss of cognition. HD is an autosomal dominantly inherited disorder and although prevalence rates range widely depending on geography and ethnicity, it is thought to affect more than 50,000 people in the United States and Europe alone. The underlying genetic cause is a trinucleotide CAG expansion in the
 25 huntingtin gene (HTT), discovered as a genetic marker by James Gusella from Massachusetts General Hospital in 1983, which results in the production of a mutant huntingtin protein (mHTT) with a toxic poly-glutamine (polyQ) tract. However, despite decades of research and clinical trials, no successful therapy has yet been developed. The "typical" onset of HD is between the ages of 40-50, but up to 15% of cases have very late onset and don't show clinical symptoms until after the age of 60. In a recent
 30 meta-analysis of studies investigating cases of late-onset HD (LoHD), more than 90% of patients had CAG repeat lengths of ≤44. One of the more interesting observations in HD is that while there is a well-known correlation between the length of the polyQ repeat tract and the onset and severity of the disease, there is substantial variability within individual patients. For example, in patients with mid-range repeat lengths (defined here as between 40 and 50), disease onset can vary by 60 years in any individual
 35 patient. This means that many patients who are carriers of polyQ tracts that predispose to the

development of the disease can live for decades in a “presymptomatic” state. What controls the onset of clinical symptoms remains currently unknown and complicates the prognostic evaluation of HD patients.

5 Although historically considered a monogenic disease, extensive research into the underlying pathology of HD suggests that the mechanisms leading to disease onset and progression are more complex than originally thought. Many different technologies have been used to look at the molecular changes underlying disease progression in HD including gene expression, proteomics, metabolomics, network analysis, genomics and single nucleotide polymorphism (SNP) profiling). As HD is considered a paradigm

10 of a disease characterized by epigenetic dysregulation, more recently epigenetic approaches have emerged as a promising new tool for assessing pathology-related changes. Most epigenetic studies in HD have focused on looking at genome wide histone modifications (acetylation, methylation) or histone modifications at specific loci related to HD. While these approaches have provided interesting insight into the disease, they have yielded often conflicting results and shown inconsistencies between mouse

15 models and human disease. As such, a consensus picture of epigenetic deregulation in HD using histone modification readouts has yet to materialize. However, not all molecular mechanisms associated with epigenetic regulation have been assessed in the context of HD. An important aspect of epigenetic regulation is at the level of 3-dimensional (3D) genomic architecture.

20 The 3D organization of the genome reflects the heterogeneous effects of external environmental cues and inputs and can be empirically measured by the assessment of chromosome conformations or when several conformations are measured concomitantly, a chromosome conformation signature (CCS). CCSs can be thought of as the molecular barcode that gives a readout of the epigenetic landscape of a given cellular population. Given the central role of mHTT in the development of HD, we hypothesized that

25 regulatory differences in genomic architecture at the HTT locus may exist between diseased individuals and healthy, unaffected controls. We used *EpiSwitch*, an established proprietary industrial platform for monitoring CCSs, to assess chromatin architecture differences between presymptomatic and symptomatic HD patients and healthy, unaffected individuals. *EpiSwitch* readouts provide high resolution, reliable and high throughput detection of CCSs while simultaneously meeting the high bar of

30 industry standards for quality control.

Methods

Sample collection

All samples were obtained from National BioService, LLC. In total, 20 samples were used in this study; 10

35 healthy control (HC) samples (CAG repeats, n<35), and 10 HD samples (CAG repeats, n>39). For the HD samples, 7 were from symptomatic patients (HD-Sym) and 3 were from presymptomatic patients who

had a diagnosis of HD but did not yet show any clinical symptoms (HD-Pre). One HD patient was taking tetrabenazine and one patient was taking sertraline. All samples were negative for *human immunodeficiency virus*, *hepatitis B virus*, *hepatitis C virus* and *syphilis* (Table 14).

5 **Study design**

We wanted to identify chromosome conformations that differed between healthy controls (low CAG), presymptomatic HD patients (high CAG, no disease manifestation) and symptomatic HD patients (high CAG, disease manifestation). We focused on a ~225 kb region surrounding the HTT locus from (chr4: 3,033,588 to 3,258,170 as annotated in hg38) for our analysis. Using the CAG repeat expansion tract in 10 exon 1 of HTT (chr4: 3,054,162 to 3,095,930) as the anchor point (“Anchor”), we defined five genomic zones surrounding the Anchor to look at chromosome conformations that varied between sample groups (Figure 2 and Table 15). These Zones were chosen based on: the presence of potential *EpiSwitch* anchoring sites; the presence of known disease related-SNPs (HD and other diseases); and the enrichment of known histone modification sites (H3K4me3, H3K36me3, and H3K27ac) in HD as found in 15 the GWASdV2 database (<http://jjwanglab.org/gwasdb>) (Figure 2).

Chromosome conformation identity

A search of the NCBI: GEO database for previously reported HD epigenetic data was performed in February 2018. Peak called ChIP-seq data for H3K4me3 from 12 (6 HD and 6 control samples) post-20 mortem prefrontal cortex brain samples (bed format) was obtained (GSE68952). In addition, Bigwig tracks of ChIP-seq data for H3K27ac and H3K36me3 from HD iPSC-derived neural cell lines and control cell lines were also downloaded (GSE95342). The data tracks were loaded into the Integrative Genome Viewer (IGV) [41] alongside the *EpiSwitch* and reference sequence annotations. Both visual and programmatic (BEDtools) comparisons were performed on the HTT locus to identify the 5 Zones of 25 interest.

Appropriate software was used to identify high probability chromatin folding interactions with one “end” occurring in the Anchor zone proximal to the CAG repeats and the other in any of the 5 Zones of interest. A total of 61 interactions matched these criteria, and for practical reasons, 20 interactions were 30 selected to cover interactions between the Anchor site and all the Zones of interest. An automated primer design application was used to design oligonucleotide pairs that amplified the expected DNA sequence caused by the interaction when subjected to the chromosome conformation capture assay.

3C and PCR

35 Chromosome conformation capture and detection by PCR were performed. Chromatin with intact chromosome conformations from 50 µl of blood sample from each patient sample was extracted using

the *EpiSwitch* assay following the manufacturer's instructions (Oxford BioDynamics Plc). Quality control on all samples was done using the detection of a chromatin loop at the MMP1 locus, a historical internal control for 3C analysis. Pooled 3C libraries for each of the sample types were generated to provide a generalized population sample for each of the sample subgroups. Real-time PCR was performed with

5 SYBR green with the CFX-96 (Bio-Rad) machine to identify the interactions with differing PCR product detection patterns between the sample types. Oligonucleotides were tested on control templates to confirm that each primer set was working correctly. In line with Royal Forensic Protocol for PCR detection the final nested PCR was performed on each sample in triplicates for the follow up data on individual HD patients. This procedure permitted the detection of limited copy-number templates with

10 higher accuracy. All PCR amplified products were monitored on the LabChip® GX from Perkin Elmer, using the LabChip DNA 1K Version2 kit (Perkin Elmer) and internal DNA markers were loaded on the DNA chip according to the manufacturer's protocol using fluorescent dyes. Fluorescence was detected by laser and electropherogram read-outs translated into a simulated band on gel picture using the instrument software. The threshold of detection for the instrument was set by the manufacturer from

15 30 fluorescence units and above.

Statistical Analysis

Data analysis was performed in R (language and environment for statistical computing). This included stats and dplyr packages for t-tests and R-squared analysis & a ggplot2 package for boxplots and

20 regression plots.

Results

Patient Clinical Characteristics

HC and HD samples were age (average 36.9 for HC and 35.3 for HD) and sex matched ($\frac{1}{2}$ male and $\frac{1}{2}$ female), with the majority (70%) of HD cases being symptomatic (**Table 16**). All samples were from Non-Hispanic or Latino Whites. Average CAG repeats lengths were 25.7 for HC and 44.2 for HD (**Table 16**). The HC and HD samples showed no statistical difference in age. The HC patients showed no statistical difference in age from HD-Sym. HD-Pre patients were younger (average age = 25.3) than HD-Sym patients (average age = 39.6) ($p = 0.02$). There was a moderate negative relationship between disease

25 duration and CAG repeat size and a moderate positive relationship between age at diagnosis and CAG repeat size, though neither were statistically significant.

There was a statistically significant increase in CAG repeat length in HD patients relative to HC ($p = 1.08$

35 E^{-7}). There was a statistically significant increase in CAG repeat length between HC and HD-Pre ($p = 3.43$

E^{-6}) and HD-Sym ($p = 9.50 E^{-8}$). There was no statistical difference in CAG repeat length between HD-Pre and HD-Sym ($p = 0.09$).

5 The average age at diagnosis for HD samples was 35.3 and the average disease duration was 3.8 years with 7 out of 10 patients reporting symptoms of irritability, chorea, or both (Table 16).

Chromosome Conformations in HC, HD-Pre and HD-Sym

10 Of the 20 interactions that were evaluated (Table 17), we identified nine informative interactions. We identified two constitutive interactions and seven conditional interactions which were present in HD, but not healthy controls. Three of the seven conditional interactions were present only in HD-Sym, and absent in HD-Pre.

Constitutive Conformations

15 All samples passed internal QC analysis for the MMP1 interaction. Two constitutive (identified in all samples) chromatin loops were identified. Both loops were between the Anchor and Zone 2 with the first loop spanning 28 kb and the second loop spanning 34 kb. Two constitutive interactions occurring in all patients (HD-Sym, HD-Pre and HC) were observed in this study. Both interactions (CC1 & CC2) were between the Anchor and Zone 2 with CC1 spanning 28 kb and CC2 spanning 34 kb.

20

Conditional Conformations

We identified seven conditional chromosome conformations that could discriminate between the different patient subgroups evaluated in this study. Specifically, we identified two chromosome interactions that were present in HC, but absent in all HD samples. The first interaction (I1) spanned the

25 Anchor and Zone 4 and covered 77 kb while the second interaction (I2) spanned the Anchor and Zone 3 and covered 140 kb. We also identified two chromosome interactions that were present in HC and HD-Pre, but absent in HD-Sym samples. Both interactions (I3 and I4) spanned the Anchor and Zone 4 and covered 92 kb and 104 kb, respectively. Last, we identified three chromosome interactions that were present in HD-Sym samples, but absent in HD-Pre and HC samples. The first interaction (I5) spanned the

30 Anchor and Zone 3, covering 122kb. Interestingly, this interaction included a SNP (rs362331) known to be a factor in the predisposition to develop HD. The second and third interactions (I6 and I7) spanned the Anchor and Zone 1 and covered 185 kb and 174 kb, respectively. Last, we tested the absence or presence of all conditional interactions in individual HD samples. In the HD-Sym samples, we found the presence of at least one of the conditional markers (I5, I6 and I7) in six out of seven samples (Figure 3).

35 A summary of all the interactions that were evaluated in this study are shown in Figure 4 and Figure 5.

Table 18 gives odds ratios. Table 13 shows chromosome interactions that did not show any differences between subgroups.

Discussion

5 **Problem Statement and Results Summary**

While it is well-known that individuals with greater than 39 CAG-repeats will get HD, the clinical onset of disease varies widely amongst individual patients and the factors that influence when the disease manifests clinically are less well characterized. Here we used *EpiSwitch*, an industrial platform for assessing chromatin architecture, to evaluate the epigenomic landscape of the HTT locus in HD patients 10 and healthy, unaffected controls. We identified a set of seven interactions that when taken together as a CCS, could differentiate HD from unaffected controls and more importantly, could differentiate between presymptomatic and symptomatic HD patients. One of these interactions, specific for symptomatic HD, contains a SNP (rs362331) shown to be associated with a predisposing disease haplogroup. When taken together, these results show that a simple, non-invasive blood-based test 15 evaluating a CCS can serve as a surrogate biomarker for assessing disease progression in HD.

Biological Relevance

While it is known that the poly-Q repeat tract expansion and production of mHTT are the underlying causes of HD, the molecular events leading to the development of clinical symptoms are less well 20 characterized. Several studies have looked at SNPs within the HTT locus as a potential contributor to disease onset. One recent SNP genotyping study of HD patients identified ~41 SNPs heterozygous in at least 30% of the patients, including the rs362331 C/T SNP in exon 50 of the HTT gene. Perhaps more biologically relevant is that when the rs362331 SNP is allele-selectively knocked down using anti-sense oligonucleotides, siRNAs or miRNA, a dramatic reduction in the levels of mHTT protein is achieved both 25 *in vitro* and *in vivo* suggesting that this SNP and its surrounding genomic landscape play an important role in regulating mHTT levels. In this study, we observed a chromosome conformation (I5) that was present only in HD-Sym patients and absent in HD-Asy and HCs that overlapped with the rs362331 SNP. This suggests that the production of neurotoxic mHTT in patients that have increased poly-Q tracts and a genetic predisposition to the early development of HD by the presence of the rs362331 SNP may be 30 regulated at the level of higher order chromatin structure. Another outstanding question in HD is how the disease is inherited in cases where neither parent has received a diagnosis. The two main prevailing hypotheses posit that 1) the carrier parent could have passed away from another factor before the onset of the disease and 2) “unstable” CAG repeat tracts expand with each generation. A third possibility also exists, in that at mid-range (35-50) repeats, individuals could be carriers without 35 manifestation of the disease, but their progeny might be unable to compensate for the genetic defect through undefined mechanisms and will develop the disease. The HD patients evaluated in this study all

had CAG repeats in this mid-range, raising the possibility that potential compensatory mechanisms in disease development may be mediated through differences in genomic architecture.

Clinical Relevance

5 HD is unique in that there exists a simple test to definitively diagnose the disease, HTT gene sequencing and measurement of CAG repeat number. For clinical care and clinical trials, there are also several tests to measure disease severity, such as the Unified Huntington's Disease Rating Scale (UHDRS), the Shoulson–Fahn Scale, and the Mini–Mental State Examination (MMSE). While these assessments measure different elements of an HD patients physical and mental well-being as a surrogate for disease 10 severity, they are all subjective in nature and most are not specific for HD. What is missing are concrete molecular tools to monitor disease progression.

There are presently 22 therapeutic agents for treating HD in different stages of preclinical and clinical development, half of which are in Phase 2 or Phase 3. Once further validated, the CCS reported here 15 could be used in clinical trials as a surrogate outcome biomarker to assess the therapeutic efficacy of the drug in question. In addition to monitoring a symptomatic patient's response to a particular therapy in clinical trials, another advantage of the approach described here lies in the information that can be obtained for presymptomatic patients. For most HD patients, the presymptomatic period can last decades. Five of the seven (i3-i7) interactions identified here clearly separate presymptomatic HD 20 patients from symptomatic ones, and when further validated could serve as an "early warning" indicator test for the onset of HD symptoms in presymptomatic carriers.

This study gives first evidence of detectable conditional differences in chromatin architecture specific for the manifestation of HD and correlated with known disease haplotypes. The major strength of this study 25 lies in the unique approach, which is based on the latest developments in understanding the regulatory role of genomic architecture. While there have been several historical studies in HD aimed at developing disease progression biomarkers based on clinical, imaging and molecular measures, to the best of our knowledge this is the first time that the assessment of higher order chromatin structures in a clinically accessible biofluid has been applied in HD.

Probe	GeneLocus	Probe_Count_Total	Probe_Count_Sig
C6orf58_6_127480771_127483471_127600017_127604343_FR	C6orf58	10	5
NEFH_22_29442588_29445314_29482081_29484217_RR	NEFH	16	10
IL1A_2_112765786_112772711_112810765_112813086_RR	IL1A	23	15
FARP1_13_98271575_98282700_98346930_98348486_FR	FARP1	26	26
PRKCA_17_66441276_66447067_66475597_66481312_RF	PRKCA	30	16
RNU6-1264P_17_6162286_6163870_6195952_6199184_FR	RNU6-1264P	31	16
PON2_7_95405100_95420940_95465337_95474032_FR	PON2	40	19
CAPN9_1_230738572_230739927_230752057_230757333_RR	CAPN9	50	23
ATXN7L1_7_105654123_105657510_105741521_105750599_FR	ATXN7L1	130	61
CNTNAP2_7_146728706_146734820_146785878_146792823_RF	CNTNAP2	144	60
CTNNA3_10_66299269_66302507_66496211_66513003_FF	CTNNA3	151	46
ZFPM2_8_105632010_105638904_105814873_105824107_FR	ZFPM2	186	68
ALDH1A2_15_58325151_58334051_58485549_58488054_FR	ALDH1A2	185	70
ALDH1A2_15_58325151_58334051_58538695_58540885_FF	ALDH1A2	185	70
MAGI2_7_79009346_79018304_79275810_79284623_RF	MAGI2	186	80
FER1L6_8_123963222_123969450_124085753_124093275_FR	FER1L6	98	13
UBQLN2_X_56536168_56538402_56570114_56575112_FR	UBQLN2	10	7

Table 1a

Table 1b

HyperG_Stats	FDR_HyperG	Percent_Sig	logFC	AveExpr	t
0.192504768	0.761792372	50	-0.206940905	-0.206940905	-7.657670128
0.012576275	0.174966022	62.5	0.155885782	0.155885782	12.07017614
0.001213793	0.03415703	65.22	0.130559454	0.130559454	3.67899153
0.0000000000000171	0.0000000000079	100	0.254822215	0.254822215	14.73928749
0.013706774	0.176284348	53.33	0.24251793	0.24251793	7.451573283
0.019964247	0.210078331	51.61	-0.152642021	-0.152642021	-9.03898498
0.032185744	0.276690843	47.5	-0.178366996	-0.178366996	-5.268810519
0.029727242	0.269876723	46	-0.224588757	-0.224588757	-8.238208751
0.000354202	0.018221718	46.92	0.189559955	0.189559955	6.597539924
0.011284771	0.170730832	41.67	-0.440614155156995	-0.440614155156995	-7.4296031698139
0.713857099	1	30.46	-0.436584531378388	-0.436584531378387	-14.2285764776972
0.122432344	0.622894915	36.56	-0.425339105073479	-0.425339105073479	-10.493629902935
0.063739285	0.421589843	37.84	0.34586199024656	0.34586199024656	13.4772416929847
0.063739285	0.421589843	37.84	0.339861526563037	0.339861526563037	7.22282802700005
0.00139007	0.035755702	43.01	0.34944598213495	0.34944598213495	10.5772811324938
0.999996196	1	13.27	0.350542987884941	0.350542987884941	5.66944350076527
0.016370015	0.18948292	70	0.353892612578078	0.353892612578078	2.69374295077173

5 Table 1c

P.Value	adj.P.Val	B	FC	FC_1	LS	Loop detected
0.000000338	8.58E-06	6.769090112	0.866372344	-1.154238137	-1	Control
0.000000000262	7.50E-08	13.89074006	1.114105444	1.114105444	1	ALS
0.001613314	0.006000711	-1.70129901	1.094718133	1.094718133	1	ALS
0.0000000000858	1.48E-08	17.19553875	1.193188696	1.193188696	1	ALS

0.0000005	1.15E-05	6.37489208	1.183055643	1.183055643	1	ALS
0.0000000281	1.46E-06	9.26023757	0.899601504	-1.111603299	-1	Control
0.000045	0.000343598	1.852676493	0.883702704	-1.131602286	-1	Control
0.000000115	3.91E-06	7.847763841	0.85583895	-1.168444134	-1	Control
0.00000027	3.98E-05	4.678642967	1.140415818	1.140415818	1	ALS
5.21452107638594E-07	1.18E-05	6.332521031	0.736820877	-1.357181957	-1	Control
1.58303956750015E-11	1.70E-08	16.61028384	0.738881785	-1.353396471	-1	Control
2.63975825723201E-09	2.70E-07	11.61504783	0.744663678	-1.342888111	-1	Control
4.03123106207005E-11	2.85E-08	15.71064142	1.270910107	1.270910107	1	ALS
7.77879601584934E-07	1.61E-05	5.930445637	1.265635109	1.265635109	1	ALS
2.32055637317047E-09	2.48E-07	11.7426843	1.27407127	1.27407127	1	ALS
0.0000188398197223922	0.000175073	2.726108027	1.275040425	1.275040425	1	ALS
0.0144551346098748	0.035577015	-3.813817915	1.278004231	1.278004231	1	ALS

Table 1d

Probe sequence		Probe Location	
60 mer		Chr	Start1
TCACCACACATCACCCCTTGCTCCTCGAGCTGGTGACCACAAACAGGGTGCCACC		6	127483440
GAGGTGGGTGAATCATGAGGTCAAGGGTTCGACAATAGTTGAGAATCTCCAACCACCTGG		22	29442590
GGCCTTATAGTCAGCTGATCAGGTGAAATCGATTGGCCTTAGGATCAGCTACCATTGC		2	112765788
GAGGCAGGCAGGATCACAAAGTCAAAAGATCGATAACTCAATAATAGTTACAGATGCAAA		13	98282669
AGCACCATATCTGGGATGTAGCTATTGCTCGAGATTGCACTGAGCTGTGATCACACCTCT		17	66441278
TCTTCCCTCTTTAAAACCACCATTCATCGACCCCCACACATCCTGTGCCACTCTACTGC		17	6163839
TAACCATTATGCATCACTAACATAGCATTGATATGATATGCTCAGTTAGTTAGGGAAA		7	95420909
GGCTCAGGAAGAGAACTATTGCTCTTCGACACGACATGCGAGACACTCACACGTAG		1	230738574
GTTGGGTGGATCCCTTGAGCTCAGGAATTGAGAATGATTTTCAGCCCCGTGGAAGG		7	105657479
ATCAAAAGAAAATAGATACTTGTCTACTCGAGTTGAATAAAATCCTCAGCTTCTGTCC		7	146728708
AAAAGAAAATGTGAAAAGTTGTCACATTCGATTAATCCAAAAGGTCTTCTATGAGGC		10	66302476
TTAAAAGTATAGTAGTTGGCATTAAACATTGACCTTTCTGTTCACTGAAACCAACCCAG		8	105638873
CATCAACTAATAGTAAACATTATAATATCGACTGAAGACCTTCATACTGTAAGATTCA		15	58334020
CATCAACTAATAGTAAACATTATAATATCGACTGCTGAGCTGAGATCACACTGCC		15	58334020
TTATTCCTTCCAAATAGTAAATTATTCGAAACTTTAAGAATCAATATAAAATTCC		7	79009348
CATAATTATAAATTAAAAATGACACTATCGATTATGTCAGTGTTCAGTTGGTGTGTC		8	123969419
CAGAGCACTAAGATAGACTTCTAAGGTTCGAGGCATAGCTCCAGCTGTATTGAGGTA	X	56538371	

Table 1e

Probe Location			4 kb Sequence Location				
End1	Start2	End2	Chr	Start1	End1	Start2	End2
127483469	127600019	127600048	6	127479470	127483469	127600019	127604018
29442619	29482083	29482112	22	29442590	29446589	29482083	29486082
112765817	112810767	112810796	2	112765788	112769787	112810767	112814766
98282698	98346932	98346961	13	98278699	98282698	98346932	98350931
66441307	66481281	66481310	17	66441278	66445277	66477311	66481310
6163868	6195954	6195983	17	6159869	6163868	6195954	6199953
95420938	95465339	95465368	7	95416939	95420938	95465339	95469338
230738603	230752059	230752088	1	230738574	230742573	230752059	230756058
105657508	105741523	105741552	7	105653509	105657508	105741523	105745522
146728737	146792792	146792821	7	146728708	146732707	146788822	146792821
66302505	66512972	66513001	10	66298506	66302505	66509002	66513001
105638902	105814875	105814904	8	105634903	105638902	105814875	105818874

58334049	58485551	58485580	15	58330050	58334049	58485551	58489550
58334049	58540854	58540883	15	58330050	58334049	58536884	58540883
79009377	79284592	79284621	7	79009348	79013347	79280622	79284621
123969448	124085755	124085784	8	123965449	123969448	124085755	124089754
56538400	56570116	56570145	X	56534401	56538400	56570116	56574115

Table 2a

probe	PCR-Primer1_ID	PCR_Primer1
C6orf58_6_127480771_127483471_127600017_127604343_FR	OBD112-021	AAGCACTTCATTCTCCCTCACC
NEFH_22_29442588_29445314_29482081_29484217_RR	OBD112-057	ACTGAGCAATGATGGCAACAC
IL1A_2_112765786_112772711_112810765_112813086_RR	OBD112-077	CTAGGCCTGCGTTCTCCGT
FARP1_13_98271575_98282700_98346930_98348486_FR	OBD112-093	AGTTCTCTCTAAGAACTCAAGGA
PRKCA_17_66441276_66447067_66475597_66481312_RF	OBD112-101	CCCCAGGCACTCACACCTTA
RNU6-1264P_17_6162286_6163870_6195952_6199184_FR	OBD112-105	GGGCACTAACACCCCTTTGT
PON2_7_95405100_95420940_95465337_95474032_FR	OBD112-125	GATGGGAATCAAGGGCAAGGG
CAPN9_1_230738572_230739927_230752057_230757333_RR	OBD112-145	GCATTAGCCAGCAAGCATAACCT
ATXN7L1_7_105654123_105657510_105741521_105750599_FR	OBD112-177	CACCGCCTGATGCCAGGTCTT
CNTNAP2_7_146728706_146734820_146785878_146792823_RF	OBD112-217	GGCACTGTTGGTCTGAAGCAC
CTNNA3_10_66299269_66302507_66496211_66513003_FF	OBD112-229	GGTCTAGATGTCAGTCAGTC
ZFPM2_8_105632010_105638904_105814873_105824107_FR	OBD112-245	GACTATAAACTCTCCTTGTCA
ALDH1A2_15_58325151_58334051_58485549_58488054_FR	OBD112-317	CCTTACCGACACCAGGTAGC
ALDH1A2_15_58325151_58334051_58538695_58540885_FF	OBD112-309	CCCGACACCAGGTAGCATT
MAGI2_7_79009346_79018304_79275810_79284623_RF	OBD112-329	GGGCACCCCCTAGACAC
FER1L6_8_123963222_123969450_124085753_124093275_FR	OBD112-333	ATTCTCTCCCTGGTAAATCCTGGT
UBQLN2_X_56536168_56538402_56570114_56575112_FR	OBD112-337	CGCCAGCTCAGCAGCAATAA

5 Table 2b

PCR-Primer2_ID	PCR_Primer2	GLMNET
OBD112-023	ATACTCCCATCCCTAGGCC	0.799327478
OBD112-059	CCCTACGACTGGCAACCCA	-0.323299991
OBD112-079	CCCTGGCATTACATCACCGA	0.422829791
OBD112-095	GTCAGCAACTGTGTCAGGG	0.471919696
OBD112-103	GGATCCACGATCTCCCTCCAC	-0.379552353
OBD112-107	AGGTCAGGATGGGTACCGTT	0.259971774
OBD112-127	CTGGGATGATTCTCTGGACTTCT	-1.132792925
OBD112-147	GGTGGGCCTGGGTTAGATGC	0.980123681
OBD112-179	CAGCTGGCCGATCCATCAC	-0.201292203
OBD112-219	GAGAACGACGACCTGGCACT	-0.613932219
OBD112-231	TCATCCTATCCTCTCCTAGC	-0.36233865
OBD112-247	ACTGTAGGCCAACAGAAAG	0.541529302
OBD112-319	CTGGTCCAGTGTCAAGCGTGT	-0.562433398
OBD112-311	GTGACTCTGCACGCACTGTT	0.486039187
OBD112-331	AGGCTCAGCAGGTTCTGCC	-0.759217304
OBD112-335	CTGGGCAGGTCACTCAGACAG	-1.154344187
OBD112-339	TGCTAGGGCCGAGTAATCATC	0.146973789

Table 3

Marker	GLMNET
OBD112-021/023	0.799327478
OBD112-057/059	-0.323299991
OBD112-077/079	0.422829791
OBD112-093/095	0.471919696
OBD112-101/103	-0.379552353
OBD112-105/107	0.259971774
OBD112-125/127	-1.132792925
OBD112-145/147	0.980123681
OBD112-177/179	-0.201292203
OBD112-217/219	-0.613932219
OBD112-229/231	-0.36233865
OBD112-245/247	0.541529302
OBD112-317/319	-0.562433398
OBD112-309/311	0.486039187
OBD112-329/331	-0.759217304
OBD112-333/335	-1.154344187
OBD112-337/339	0.146973789

Table 4a

Genes	Full Name
C6orf58	Chromosome 6 Open Reading Frame 58
NEFH	Neurofilament Heavy
IL1A	Interleukin 1 Alpha
FARP1	FERM, ARH/RhoGEF And Pleckstrin Domain Protein 1
PRKCA	Protein Kinase C Alpha
RNU6-1264P	RNA, U6 Small Nuclear 1264, Pseudogene
PON2	Paraoxonase 2
CAPN9	Calpain 9
ATXN7L1	Ataxin 7 Like 1
CNTNAP2	Contactin Associated Protein-Like 2
CTNNA3	Catenin Alpha 3
ZFPM2	Zinc Finger Protein, FOG Family Member 2
ALDH1A2	Aldehyde Dehydrogenase 1 Family Member A2
MAGI2	Membrane Associated Guanylate Kinase, WW And PDZ Domain Containing 2
FER1L6	Fer-1 Like Family Member 6
UBQLN2	Ubiquilin 2

5 Table 4b

Activity
Protein Coding gene
maintenance of neuronal caliber, intracellular transport to axons and dendrites, neuronal damage and susceptibility to amyotrophic lateral sclerosis (ALS)
various immune responses, inflammatory processes, hematopoiesis and cell injury, activates apoptosis, correlated with rheumatoid arthritis and Alzheimer's disease
dendritic growth in neurons, link between cytoskeleton to the cell membrane
diverse cellular signalling pathways, receptors for phorbol esters, cellular processes such as cell adhesion, cell transformation, cell cycle checkpoint, and cell volume control

Pseudogene, connected with the snRNA class
cellular antioxidant, human tissues and membrane-bound, defence responses to pathogenic bacteria, protecting cells from oxidative stress
neurodegenerative processes and digestive tract, associated with gastric cancer
Protein Coding gene
regulates interactions between neurons and glia, associated with speech and language development, cell adhesion in nervous system
cell-cell adhesion in muscle cells, related to arrhythmogenic right ventricular dysplasia
regulates hematopoiesis and cardiogenesis in mammals,
developing and adult tissues, posterior organ development, prevents spina bifida
dentatorubral and pallidoluysian atrophy
Protein Coding gene
in vivo protein degradation

Table 5a

Array_data_for_3_month_Samples

probe	GeneLocus	Probe_Count_Total	Probe_Count_Sig
ZFPM2_8_104964347_104973135_105209447_105218050_FF	ZFPM2	187	6
VEGFA_6_43733863_43737741_43777557_43780533_FR	VEGFA	42	1
ERBB4_2_212287088_212294815_212317329_212325591_FR	ERBB4	193	10
ZFPM2_8_105209447_105218050_105302264_105310138_FF	ZFPM2	187	6
PASD1_X_151600201_151608969_151687203_151692271_FR	PASD1	41	14
GRM7_3_7259138_7267165_7394377_7401227_RF	GRM7	174	7
GRIK2_6_102047545_102055341_102078831_102089392_FR	GRIK2	188	5
SORCS2_4_7219417_7226457_7241158_7248673_FF	SORCS2	84	6
LINGO2_9_28314155_28333777_28371887_28374860_FR	LINGO2	194	3

5

Table 5b

HyperG_Stats	FDR_HyperG	Percent_Sig	logFC	AveExpr	t	P.Value	adj.P.Val
0.067854625	0.99237389	3.21	0.272052749	7.772607282	3.665534368	0.001024476	0.999953189
0.475978534	1	2.38	-0.142175177	14.62826333	-2.054497284	0.049386319	0.999953189
0.000818098	0.03828697	5.18	0.383778852	7.597586859	3.176083618	0.003622402	0.999953189
0.067854625	0.99237389	3.21	0.229570464	9.577318093	2.801385891	0.00913346	0.999953189
7.57E-16	3.54E-13	34.15	0.341166309	9.148011765	2.62030276	0.01404442	0.999953189
0.01784403	0.37959119	4.02	0.306594489	9.301524803	2.334933566	0.026957376	0.999953189
0.16129386	1	2.66	0.286167252	9.345860096	2.824849044	0.008630812	0.999953189
0.001827031	0.0712542	7.14	-0.222042969	10.03535881	-2.171963051	0.038499943	0.999953189
0.570146657	1	1.55	0.40947654	6.9839738	2.811551759	0.008912387	0.999953189

Table 5c

B	FC	FC_1	LS	Loop detected
-1.464031868	1.207524741	1.207524741	1	Fs
-3.802493578	0.906151904	-1.103567731	-1	Sw
-2.223740309	1.304754929	1.304754929	1	Fs
-2.785028975	1.172485811	1.172485811	1	Fs
-3.046315984	1.266780276	1.266780276	1	Fs
-3.440610909	1.236784796	1.236784796	1	Fs

-2.750635647	1.219396451	1.219396451	1	Fs
-3.654353119	0.857350501	-1.166384109	-1	Sw
-2.770141569	1.328203808	1.328203808	1	Fs

5 Table 5d

Probe sequence 60 mer	Probe Location		
	Chr	Start1	End1
TATATTAAAAATACATACTGGTATACATCGATCTCATGACTTTGCTATTATGCATAGTG	8	104973104	104973133
CCCCAGCCAGCAACCTGGCTCACCTGATCGAGTACATCTCAAGCCATCTGTGTGCC	6	43737710	43737739
TATAAAATAATACAGCTCTATTGCCTACTCGATTAAGAATCATATTATATCCTTAATTC	2	212294784	212294813
CACTATGCATAATAGCAAAGTCATGAGATCGAAAATGTTGTCAAGCAGTAGGTTTGGG	8	105218019	105218048
GATTTAGAATCTCTAACAAAGGCTGCAATCGAGGTTAGCTGCTGCAGAAAGAAGAGAAA	X	151608938	151608967
GGTGACTAAATGAGATTGCATTTCTTCGACCATTGGCCAGCATGCCAAACACTGGT	3	7259140	7259169
TAACCTCTCTTCTTAGGTTCTCCATATCGATAGAAAATTGCTGCAGCCCTTAATGCC	6	102055310	102055339
AGCTCACGGTCAGTGCGTCCGTTGCTCGAATAAGAACAGGACCTAAAAATAGA	4	7226426	7226455
AAAGTCATACAACACTATGTAAGATATCGAACACTGTTAGAATAGGTGAAGGTTAT	9	28333746	28333775

Table 5e

Probe Location Start2	End2	4 kb Sequence Location				
		Chr	Start1	End1	Start2	End2
105218019	105218048	8	104969134	104973133	105214049	105218048
43777559	43777588	6	43733740	43737739	43777559	43781558
212317331	212317360	2	212290814	212294813	212317331	212321330
105310107	105310136	8	105214049	105218048	105306137	105310136
151687205	151687234	X	151604968	151608967	151687205	151691204
7401196	7401225	3	7259140	7263139	7397226	7401225
102078833	102078862	6	102051340	102055339	102078833	102082832
7248642	7248671	4	7222456	7226455	7244672	7248671
28371889	28371918	9	28329776	28333775	28371889	28375888

Table 6a

10 Array_data_for_6_month_Samples

probe	GeneLocus	Probe_Count_Total	Probe_Count_Sig	HyperG_Stats
ZFPM2_8_104964347_104973135_105209447_105218050_FF	ZFPM2	187	8	0.007868339
VEGFA_6_43733863_43737741_43777557_43780533_FR	VEGFA	42	1	0.472787206
ERBB4_2_212287088_212294815_212317329_212325591_FR	ERBB4	193	13	8.61E-06
ZFPM2_8_105209447_105218050_105302264_105310138_FF	ZFPM2	187	8	0.007868339
PASD1_X_151600201_151608969_151687203_151692271_FR	PASD1	41	18	1.87E-22
GRM7_3_7259138_7267165_7394377_7401227_RF	GRM7	174	6	0.049498361
GRIK2_6_102047545_102055341_102078831_102089392_FR	GRIK2	188	6	0.066833261
SORCS2_4_7219417_7226457_7241158_7248673_FF	SORCS2	84	3	0.134527864
LINGO2_9_28314155_28333777_28371887_28374860_FR	LINGO2	194	4	0.337532174

Table 6b

FDR_HyperG	Percent_Sig	logFC	AveExpr	t	P.Value	adj.P.Val
0.204139683	4.28	0.214103447	7.772607282	2.884747708	0.007463232	0.999935724

1	2.38	-0.142760759	14.62826333	-2.062959216	0.04851942	0.999935724
0.001005104	6.74	0.365781024	7.597586859	3.027136877	0.005258488	0.999935724
0.204139683	4.28	0.23990173	9.577318093	2.927455517	0.006723676	0.999935724
8.73E-20	43.9	0.331193838	9.148011765	2.543709927	0.016785386	0.999935724
0.825561956	3.45	0.354666043	9.301524803	2.701032396	0.011610123	0.999935724
1	3.19	0.294782642	9.345860096	2.909894328	0.007018965	0.999935724
1	3.57	-0.229277936	10.03535881	-2.242733507	0.033029594	0.999935724
1	2.06	0.440106238	6.9839738	3.021861688	0.005327738	0.999935724

Table 6c

B	FC	FC_1	LS	Loop detected
-2.662323695	1.159982827	1.159982827	1	Fs
-3.791998096	0.905784176	-1.104015754	-1	Sw
-2.449689974	1.288579031	1.288579031	1	Fs
-2.598930246	1.18091222	1.18091222	1	Fs
-3.154405805	1.25805399	1.25805399	1	Fs
-2.930776901	1.278689554	1.278689554	1	Fs
-2.625038885	1.226700147	1.226700147	1	Fs
-3.562656794	0.853061739	-1.172248096	-1	Sw
-2.457630227	1.35670423	1.35670423	1	Fs

Fs=Faster Sw=Slower

5 Table 6d

Probe sequence 60 mer	Probe Location		
	Chr	Start1	End1
TATATTTAAAAATACATACTGGTATACATCGATCTCATGACTTGTCTATTATGCATAGTG	8	104973104	104973133
CCCCAGCCCAGCAACCTGGCTCACCTGATCGAGTACATCTCAAGCCATCCTGTGTGCC	6	43737710	43737739
TATAAATAATACAGCTCTTTGCCTACTCGATTAAAGAACATATTATATCCTTAATT	2	212294784	212294813
CACTATGCATAATAGCAAAGTCATGAGATCGAAAATGTTGTCAAGCAGTAGGTTGGG	8	105218019	105218048
GATTTAGAACATCTAACAAAGGCTGCAATCGAGGTTAGCTGCTGCAGAAAGAACAGAGAAA	X	151608938	151608967
GGTGAATAAATGAGATTGCATTTCTTCGACCATTGGCCAGCATGCCAACACTGGT	3	7259140	7259169
TAACCTCTCCCTTCTAGTTCTCCATATCGATAGAAAATTGTCTGCAGCCCTTAATGCC	6	102055310	102055339
AGCTCACGGTCAGTGCCGTTGCTCGAATAAGAACAGGACCTAAAAATAGA	4	7226426	7226455
AAAGTCATACAACTACTATGTAAGATATTGAATACCTGTTAGAATAGGTGAAGGTTAT	9	28333746	28333775

Table 6e

Probe Location	4 kb Sequence Location						
	Start2	End2	Chr	Start1	End1	Start2	End2
105218019	105218048	8	104969134	104973133	105214049	105218048	
43777559	43777588	6	43733740	43737739	43777559	43781558	
212317331	212317360	2	212290814	212294813	212317331	212321330	
105310107	105310136	8	105214049	105218048	105306137	105310136	
151687205	151687234	X	151604968	151608967	151687205	151691204	
7401196	7401225	3	7259140	7263139	7397226	7401225	
102078833	102078862	6	102051340	102055339	102078833	102082832	
7248642	7248671	4	7222456	7226455	7244672	7248671	
28371889	28371918	9	28329776	28333775	28371889	28375888	

Table 7a

probe	Inner_primers	
	PCR- Primer1_ID	PCR_Primer1
ZFPM2_8_104964347_104973135_105209447_105218050_FF	OBD112_773	GCAGCACACAGGGAACTCTCTT
VEGFA_6_43733863_43737741_43777557_43780533_FR	OBD112_765	GCTAGGAAGGGCCTGGGATG
ERBB4_2_212287088_212294815_212317329_212325591_FR	OBD112_477	TCCAAATGTTAATACTGCCTAGA
ZFPM2_8_105209447_105218050_105302264_105310138_FF	OBD112_777	CAGCCACTGTAGAGAGCAGT
PASD1_X_151600201_151608969_151687203_151692271_FR	OBD112_637	ACTTCTTCCAAGTCACTTTGC
GRM7_3_7259138_7267165_7394377_7401227_RF	OBD112_517	TGACGAAGAACAAATCCCTGGT
GRIK2_6_102047545_102055341_102078831_102089392_FR	OBD112_501	AATCTCTGCCCTCCTCTCATTTG
SORCS2_4_7219417_7226457_7241158_7248673_FF	OBD112_725	AGGTATGCAGCCAGCCTGAG
LINGO2_9_28314155_28333777_28371887_28374860_FR	OBD112_561	CCGTGCCATATCCTCTGATTTATGC

Table 7b

Inner_primers		
PCR-Primer2_ID	PCR_Primer2	GLMNET
OBD112_775	TTGTTGAGCCCAGCAATTCTTT	-0.18669073
OBD112_767	CTCAGTGGGCACACACTCCA	0.15278372
OBD112_479	AGCCATGTGGCTGGAATCT	-0.24015518
OBD112_779	TAACCCACCAGCAGCAAGGT	0.22911033
OBD112_647	TGGCCATCTTGCTTGCCCTC	0.52468015
OBD112_519	GGACCTACCTCCACTGGGTTG	0.10503793
OBD112_503	CATGTTCCCACAGCAAGGAAGTTA	-0.13060979
OBD112_727	TTTCCGTGCCAGTGTCTGT	-0.1217346
OBD112_563	GGCTGACCTCAACAGATT CGC	0.01500745

5 Table 8

Gene	Marker	GLMNET
ERBB4_2	OBD112_477.479_0.4ng	-0.24015518
ZFPM2_8	OBD112_773.775_6.5ng	-0.18669073
GRIK2_6	OBD112_501.503_0.8ng	-0.13060979
SORCS2_4	OBD112_725.727_3.25ng	-0.1217346
LINGO2_9	OBD112_561.563_13ng	0.01500745
GRM7_3	OBD112_517.519_13ng	0.10503793
VEGFA_6	OBD112_765.767_13ng	0.15278372
ZFPM2_8	OBD112_777.779_6.5ng	0.22911033
PASD1_X	OBD112_637.647_3.25ng	0.52468015

Table 9

Genes	Full name	Activity
ZFPM2	Zinc Finger Protein, FOG Family Member 2	transcription factor, regulates the activity of GATA family protein, regulates GATA-target genes expression
VEGFA	Vascular Endothelial Growth Factor A	stimulates proliferation and migration of vascular endothelial cells, stimulates physiological and pathological angiogenesis, modulates tumor stage and progression

ERBB4	Erb-B2 Receptor Tyrosine Kinase 4	stimulates a variety of cellular responses including mitogenesis and differentiation,
PASD1	PAS Domain Containing Repressor 1	encodes a cancer-associated antigen, induces autologous T-cell responses, immunotherapeutic target for the treatment of various hematopoietic malignancies
GRM7	Glutamate Metabotropic Receptor 7	activates ionotropic and metabotropic glutamate receptors, involved in most aspects of normal brain function, perturbed in neuropathologic conditions
GRIK2	Glutamate Ionotropic Receptor Kainate Type Subunit 2	modulates neurophysiologic processes
SORCS2	Sortilin Related VPS10 Domain Containing Receptor 2	protein coding gene, over expression in the central nervous system
LINGO2	Leucine Rich Repeat And Ig Domain Containing 2	protein coding gene

Table 10a

N	Array_Probe	Primer1_ID
1	DCLK1_13_35749195_35757694_36000600_36005542_RR	OBD112_453
2	GRM7_3_7259138_7267165_7394377_7401227_RF	OBD112_517
3	LINGO2_9_28314155_28333777_28371887_28374860_FR	OBD112_561
4	PASD1_X_151600201_151608969_151687203_151692271_FR	OBD112_637
5	UBQLN2_X_56536168_56538402_56570114_56575112_RR	OBD112_757
6	ZFPM2_8_104964347_104973135_105209447_105218050_FF	OBD112_773
7	ZFPM2_8_105209447_105218050_105302264_105310138_FF	OBD112_777
8	ZNF804B_7_89108281_89114839_89127203_89146680_RF	OBD112_797

Table 10b

Primer1_Seq	Primer2_ID	Primer2_Seq
TCTTGTACACGGTTGGTGGT	OBD112_455	TGTCACCTATGTGCTGAGTACTGG
TGACGAAGAAGCAATCCCTGGT	OBD112_519	GGACCTACCTCCACTGGGTTG
CCGTGCCATATCCTCTGATTATGC	OBD112_563	GGCTGACCTTCAACAGATTGCG
ACTTCTCCAAGTCACTTTTGC	OBD112_647	TGGCCATTTGCTTGCCTC
GGATATGCAGTTTCTGGCACTAC	OBD112_759	CATGCTAGGGCCGAGTAATCATCT
GCAGCACACAGGAACTCTCTT	OBD112_775	TTGTTGAGCCCAGCAATTCTTT
CAGCCACTGTAGAGAGCAGT	OBD112_779	TAACCCACCAGCAGCAAGGT
AGTAGCTTCCCTGTTAGAGGTCTTG	OBD112_799	AGCCAGTGACTCCACAACTTCTT

Table 10c

Hydrolysis_probe_ID	Hydrolysis_probe_Seq
OBD112_88_FAM	AGGTTGCTTCGAAGTACAGATATCACT
OBD112_57_FAM	ATTTCTTCGACCATTGGCCAGCA
OBD112_90_FAM	TTCTAACAGGTATTCGAATATCTTACATAGT
OBD112_26_FAM	AAGTAACATTGACAAACCTCTGGTAA
OBD112_95_FAM	TATTCTCTTGATCGAGGCATATAGCT
OBD112_80_FAM	AAAGTCATGAGATCGATGTATACCAAGTAT
OBD112_32_FAM	TTGACAAACATTTCGATCTCATGACTT
OBD112_33_FAM	TGTGGCTCAGTCGACAGAAAGTACA

Table 11a (i)

probe	GeneLocus	Probe_Count_Total	Probe_Count_Sig
DCLK1_13_35749195_35757694_36000600_36005542_RR	DCLK1	182	3
UBQLN2_X_56536168_56538402_56570114_56575112_RR	UBQLN2	10	7
ZNF804B_7_89108281_89114839_89127203_89146680_RF	ZNF804B	191	10

5 Table 11a (ii)

HyperG_Stats	FDR_HyperG	Percent_Sig	logFC	AveExpr	t
0.254316537	1	1.65	0.199442299	7.402197005	2.87151
0.000000000001	0.000000000386	70	0.283366605	9.419823948	2.114955
1.74E-05	0.002050325	5.24	0.228756351	9.389157739	2.52348

Table 11a (iii)

P.Value	adj.P.Val	B	FC	FC_1	LS	Loop detected
0.007707652	0.999953189	-2.681902147	1.14825439	1.14825439	0	Faster
0.043482517	0.999953189	-3.726899057	1.217031581	1.217031581	0	Faster
0.017587875	0.999953189	-3.182692934	1.171824364	1.171824364	0	Faster

Table 11b (i)

probe	PCR-Primer1_ID	PCR_Primer1
DCLK1_13_35749195_35757694_36000600_36005542_RR	OBD112_453	TCTTGTACACGGTTGGTGGT
UBQLN2_X_56536168_56538402_56570114_56575112_RR	OBD112_757	GGATATGCAGTTTCTGGCACTAC
ZNF804B_7_89108281_89114839_89127203_89146680_RF	OBD112_797	AGTAGCTCCCTGTTAGAGGTCTTG

10

Table 11b (ii)

PCR-Primer2_ID	PCR_Primer2	GLMNET
OBD112_455	TGTCACCTATGTGCTGAGTACTGG	0.047800994
OBD112_759	CATGCTAGGGCCGAGTAATCATCT	-0.013191036
OBD112_799	AGCCAGTGACTCCACAACTTCTT	-0.025426843

Table 11c

Marker	GLMNET
OBD112_453.455_3.25ng	0.047801
OBD112_757.759_13ng	-0.013191
OBD112_797.799_1.6ng	-0.025427

Table 12a

				Probe sequence	
				60 mer	
	probe	GeneLocus	Loop detected		
HTT_4_3088056_3090591_3182426_3183765_FR	HTT	HD_Negative		AGATCTAGTTCACAGTAGCACCATAATCGACAGATACTGACATCATCCTCCAAATGT	
HTT_4_3076247_3078445_3182426_3183765_RR	HTT	HD_Negative		AGAGTACTTCCCTAACCTACTGTACACTCGACAGATACTGACATCATCCTCCAAATGT	
HTT_4_3062575_3066389_3251623_3253432_RF	HTT	HD_Positive		TATAACCAAGTGCTCCCTAACCTACTGTACACTCGAGGATGATCGCTCCGACAGCTCCAGC	
HTT_4_3076247_3078445_3253432_3258164_RR	HTT	HD_Positive		AGAGTACTTCCCTAACCTACTGTACACTCGAGGATGATCGCTCCGACAGCTCCAGC	
HTT_4_3070353_3072847_3212506_3214871_FR	HTT	HD_Negative		GGGGTTTCGCCATGTTGGCCAGGCTGGTCTCGAAAGTTGATGCACTGTGCTCAGTTTGCA	
HTT_4_3088056_3090591_3212506_3214871_FF	HTT	HD_Positive		AGATCTAGTTCACAGTAGCACCATAATCGACTGTCCTGTTGGCCTATCTCACCCCT	
HTT_4_3088056_3090591_3167292_3170536_FR	HTT	HD_Negative		AGATCTAGTTCACAGTAGCACCATAATCGAACTCCCTGACCTTGATCCACCCACCTC	

Table 12b

Probe Location				probe	PCR_Primer1_ID	PCR_Primer1
Chr	Start1	End1	Start2	End2		
4	3090560	3090591	3182426	3182457	HTT_4_3088056_3090591_3182426_3183765_FR	RD031_037
4	3076247	3076278	3182426	3182457	HTT_4_3076247_3078445_3182426_3183765_RR	RD031_061
4	3062575	3062606	3253401	3253432	HTT_4_3062575_3066389_3251623_3253432_RF	RD031_185
4	3076247	3076278	3253432	3253463	HTT_4_3076247_3078445_3253432_3258164_RR	RD031_237
4	3072816	3072847	3212506	3212537	HTT_4_3070353_3072847_3212506_3214871_FR	RD031_241
4	3090560	3090591	3214840	3214871	HTT_4_3088056_3090591_3212506_3214871_FF	RD031_329
4	3090560	3090591	3167292	3167323	HTT_4_3088056_3090591_3167292_3170536_FR	RD031_333

Table 12c

PCR- Primer1_ID	PCR_Primer2
RD031_039	GATTCCAGCACCACCTTACAAG
RD031_063	TGATTCCAGCCACCCACCTTCACAA
RD031_187	GAACCGCACCCTACCTCAGCAGT
RD031_239	GGACAAGCAGACACACTACCTGAAC
RD031_243	ATCCCCCTGAACAGAAAGGACCTCGTG
RD031_331	GAGCCGCTCTCATATAAACCTCAGGGT
RD031_335	CAGTGGTTAGGGCAAAGAGAGGGAG

Table 13a

		Probe sequence	
		60 mer	
probe	GeneLocus	Loop detected	
HTT_4_3088056_3090591_3112545_3114867_FR	HTT	No difference	AGATCTAGTTACAGTAGCACAATATTCGACATTAAAGTTCATGAGAGTCCTCTATATT
HTT_4_3062575_3066389_3119060_3125597_RR	HTT	No difference	GCGCACAGGAGAAAGTGAAGTTGAGACCTCGAAAAAGGAATAAAATGAAAAATAGAGGAA
HTT_4_3076247_3078445_3241592_3247577_RR	HTT	No difference	AGAGTACTCCCTAACCTCTACTGTACACTCGATGTTGGTATTGAATGTGGTAAGTGG
HTT_4_3080953_3084487_3253432_3258164_RR	HTT	No difference	ATCAAGAGACTGTATGGTACTGGCACAGGATCGAGGATGATCGCTCCGACAGCTCCAGC
HTT_4_3080953_3084487_3241592_3247577_RR	HTT	No difference	ATCAAGAGACTGTATGGTACTGGCACAGGATCGATGTTGGTATTGAATGTGGTAAGTGG
HTT_4_3080953_3084487_3112545_3114867_RR	HTT	No difference	ATCAAGAGACTGTATGGTACTGGCACAGGATCGACATAAGTTCATGAGAGTCCTCTATATT
HTT_4_3076247_3078445_3112545_3114867_RF	HTT	No difference	TGTCCAGTTAAATTGGAGATTTCCGATCGAGTGTACAGTAGGAGTTAGGAAGTACTCT
HTT_4_3080953_3084487_3167292_3170536_RR	HTT	No difference	ATCAAGAGACTGTATGGTACTGGCACAGGATCGAAACTCTGACCTTGATCCACCCACCTC
HTT_4_3038381_3040238_3088056_3090591_RF	HTT	No difference	AGATCTAGTTACAGTAGCACAATATTCGAGGATGGTCTGAAACTCTGACCTTGATCCACCCACCTC
HTT_4_3088056_3090591_3253432_3258164_FR	HTT	No difference	GGGTTTCACCATGTTGCCAGGATGGTCTGAAACTCTGACCTTGATCCACCCACCTC
HTT_4_3062575_3066389_3167292_3170536_FR	HTT	No difference	AGATCTAGTTACAGTAGCACAATATTCGAGGATGGTCTGAAACTCTGACCTTGATCCACCCACCTC
HTT_4_3088056_3090591_3251623_3253432_FF	HTT	No difference	AGATCTAGTTACAGTAGCACAATATTCGAAAGGGCCCTGTTAGGAGCACTGGTTATA
HTT_4_3088056_3090591_3241592_3247577_FF	HTT	No difference	AGATCTAGTTACAGTAGCACAATATTCGACTAAGGTAGCCCTCGTGCTGGGGCT

Table 13b

Probe Location					PCR-Primer1_ID	PCR_Primer1
Chr	Start1	End1	Start2	End2	probe	
4	3090560	3090591	3112545	3112576	HTT_4_3088056_3090591_3112545_3114867_FR	RD031_001
4	3062575	3062606	3119060	3119091	HTT_4_3062575_3066389_3119060_3125597_RR	RD031_009
4	3076247	3076278	3241592	3241623	HTT_4_3076247_3078445_3241592_3247577_RR	RD031_025
4	3080953	3080984	3253432	3253463	HTT_4_3080953_3084487_3253432_3258164_RR	RD031_041
4	3080953	3080984	3241592	3241623	HTT_4_3080953_3084487_3241592_3247577_RR	RD031_057
4	3080953	3080984	3112545	3112576	HTT_4_3080953_3084487_3112545_3114867_RR	RD031_065
4	3076247	3076278	3114836	3114867	HTT_4_3076247_3078445_3112545_3114867_RF	RD031_097
4	3080953	3080984	31167292	31167323	HTT_4_3080953_3084487_3167292_3170536_RR	RD031_149
4	3038381	3038412	3090560	3090591	HTT_4_3038381_3040238_3088056_3090591_RF	RD031_157
4	3090560	3090591	3253432	3253463	HTT_4_3088056_3090591_3253432_3258164_FR	RD031_205
4	3066358	3066389	31167292	31167323	HTT_4_3062575_3066389_31167292_3170536_FR	RD031_261
4	3090560	3090591	3253401	3253432	HTT_4_3088056_3090591_3251623_3253432_FF	RD031_301
4	3090560	3090591	3247546	3247577	HTT_4_3088056_3090591_3241592_3247577_FF	RD031_337

Table 13c

PCR- Primer2_ID	PCR_Primer2
RD031_003	GCACTGCTCGCAATAGCCAAGAACTA
RD031_011	CTTGTGACCTCCCTATGGATGG
RD031_027	GCACAGAGTCCAACATTCCAC
RD031_043	GGACAAGCAGACACACTACCTGAAC
RD031_059	GCACAGAGTTCCAACATTCCAC
RD031_067	GCACTGCTCGCAATAGCCAAGAACTA
RD031_099	ACCAAATGCCATCTGGACACATCCA
RD031_151	ACAGTGGTTAGGGCAAAGAGGGGA
RD031_159	GAAAGAAGCCCTCTGGTCTGGAAA
RD031_207	GGAAAGGACAAGCAGACACACTACCTG
RD031_263	GTGGTTAGGGCAAAGAGGGGAGAT
RD031_303	TCCCGAGTCTGTGATGGCAAACCT
RD031_339	ATGGCTACGGAAAGGGCATTCGGAC

Table 14

Subject ID	Type of specimen	Specimen date	Age at collection	Gender	Specimen collection date	Specimen collection time	Specimen collection location	Specimen recorded date	Specimen recorded time	Specimen recorded location	Specimen test date	Specimen test time	Specimen test location	Specimen test result	Specimen test if available	Specimen test if available
control 01	Control	Whole blood 9 mL	32	F	2017-11-07	Caucasian	-	PCR	n1-17	no	no	no	neg	neg	neg	
control 02	Control	Whole blood 9 mL	65	F	2017-11-10	Caucasian	-	PCR	n1-35	no	no	no	neg	neg	neg	
control 03	Control	Whole blood 9 mL	41	M	2017-11-13	Caucasian	-	PCR	n1-35	no	no	no	neg	neg	neg	
control 04	Control	Whole blood 9 mL	19	M	2017-11-10	Caucasian	-	PCR	n1-17	no	no	no	neg	neg	neg	
control 05	Control	Whole blood 9 mL	46	M	2017-11-10	Caucasian	-	PCR	n1-15	no	no	no	neg	neg	neg	
control 06	Control	Whole blood 9 mL	29	F	2017-11-13	Caucasian	-	PCR	n1-35	no	no	no	neg	neg	neg	
control 07	Control	Whole blood 9 mL	47	F	2017-11-14	Caucasian	-	PCR	n1-35	no	no	no	neg	neg	neg	
control 08	Control	Whole blood 9 mL	22	M	2017-11-14	Caucasian	-	PCR	n1-19	no	no	no	neg	neg	neg	
control 09	Control	Whole blood 9 mL	35	F	2017-11-15	Caucasian	-	PCR	n1-35	no	no	no	neg	neg	neg	
control 11	Control	Whole blood 9 mL	33	M	2017-11-16	Caucasian	-	PCR	n2-35	no	no	no	neg	neg	neg	
HD 01	Huntington's	Whole blood 9 mL	32	M	2017-10-30	Caucasian	27	PCR	n1-35	no	no	no	neg	neg	neg	
HD 02	Huntington's	Whole blood 9 mL	32	F	2017-10-30	Caucasian	29	PCR	n1-35	no	no	no	neg	neg	neg	
HD 03	Huntington's	Whole blood 9 mL	28	F	2017-11-01	Caucasian	25	PCR	n1-20	Irritability, chorea	no	no	neg	neg	neg	
HD 12	Huntington's	Whole blood 9 mL	50	M	2017-11-02	Caucasian	46	PCR	n1-20	Irritability, chorea	no	no	neg	neg	neg	
HD 05	Huntington's	Whole blood 9 mL	52	M	2017-11-02	Caucasian	49	PCR	n1-35	Irritability, chorea	no	no	neg	neg	neg	
HD 06	Huntington's	Whole blood 9 mL	43	M	2017-11-02	Caucasian	35	PCR	n1-22	Irritability, chorea	no	no	neg	neg	neg	
HD 08	Huntington's	Whole blood 9 mL	23	F	2017-11-02	Caucasian	-	PCR	n1-35	Irritability, chorea	no	no	neg	neg	neg	
HD 09	Huntington's	Whole blood 9 mL	21	F	2017-11-02	Caucasian	-	PCR	n2-41	Irritability, chorea	no	no	neg	neg	neg	
HD 10	Huntington's	Whole blood 9 mL	38	F	2017-11-02	Caucasian	36	PCR	n1-35	Irritability, chorea	no	no	neg	neg	neg	
HD 11	Huntington's	Whole blood 9 mL	34	M	2017-11-15	Caucasian	32	PCR	n1-24	Irritability, chorea	no	no	neg	neg	neg	

Table 15

Zone	hg19 coordinates	hg19 coordinates	hg38 coordinates
Anchor point CAG repeat site	chr4:3055890-3097657		chr4:3054163-3095930
Zone 1	chr4:3232680-3259897		chr4:3230953-3258170
Zone 2	chr4:3107637-3130235		chr4:3105908-3128508
Zone 3	chr4:3212999-3218545		chr4:3211272-3216818
Zone 4	chr4:3161885-3185826		chr4:3160158-3184099
Zone 5	chr4:3033588-3050187		chr4:3031881-3048460

Table 16

	Healthy Controls (N=10)	Huntington's Disease (N=10)
Gender	Male (N, (%))	5 (50)
	Female (N, (%))	5 (50)
Ethnicity	Non-Hispanic or Latino	10 (100)
		10 (100)
Race	White (N, (%))	10 (100)
		10 (100)
Huntington's Type	Symptomatic (N, (%))	N/A
	Asymptomatic (N, (%))	N/A
CAG repeat length (Average, (SD))	25.7 (5.4)	7 (70)
		3 (30)
Age at Diagnosis (Average, (SD))	N/A	44.2 (2.6)
		34.9* (8.1)
Age at Sample Collection (Average, (SD))	36.9 (12.8)	35.3 (10.0)
		3.8 (1.9)
% Reporting Irritability	N/A	70
		50
% Reporting Chorea	N/A	

*Age at Diagnosis was not available for 2 of the 10 HD patients

N/A = Not Applicable

Table 17

Interaction ID	Zone3	Zone2	Zone1	Anchor	Zone3	Zone2	Zone1	Zone3	Zone2	Zone1
803031_325/27	/	/	/	/	/	/	/	/	/	/
803031_341/43	/	/	/	/	/	/	/	/	/	/
803031_357/58	/	/	/	/	/	/	/	/	/	/
803031_361/63	/	/	/	/	/	/	/	/	/	/
803031_385/57	/	/	/	/	/	/	/	/	/	/
803031_387/388	/	/	/	/	/	/	/	/	/	/
803031_388/111	/	/	/	/	/	/	/	/	/	/
803031_349/31	/	/	/	/	/	/	/	/	/	/
803031_237/338	/	/	/	/	/	/	/	/	/	/
803031_284/283	/	/	/	/	/	/	/	/	/	/
803031_357/459	/	/	/	/	/	/	/	/	/	/
803031_301/3	/	/	/	/	/	/	/	/	/	/
803031_328/331	/	/	/	/	/	/	/	/	/	/
803031_321/303	/	/	/	/	/	/	/	/	/	/
803031_337/338	/	/	/	/	/	/	/	/	/	/
803031_337/338	/	/	/	/	/	/	/	/	/	/
803031_185/187	/	/	/	/	/	/	/	/	/	/
803031_285/287	/	/	/	/	/	/	/	/	/	/
803031_241/313	/	/	/	/	/	/	/	/	/	/
803031_333/335	/	/	/	/	/	/	/	/	/	/

Table 18a

Marker	HD01	HD03	HD05	HD06	HD10	HD11	HD12	Detection rates
RD031_237.239	0	0	0	1	1	1	1	57.14
RD031_329.331	0	1	1	1	1	1	1	85.71
RD031_185.187	0	0	1	1	0	1	0	42.86

Table 18b

237.239	Symptoms	No Symptoms		
Loop Present	4	1	risk ratio	0.8
Loop Absent	3	12	risk ratio	4

Table 18c

329.331	Symptoms	No Symptoms		
Loop Present	6	1	risk ratio	0.85714
Loop Absent	1	12	risk ratio	11.1429

58

Table 18d

185.187	Symptoms	No Symptoms		
Loop Present	3	1	risk ratio	0.75
Loop Absent	4	12	risk ratio	3

Odds ratio

9

Table 19

>GRN07_3_7259138_7267165_7394377_7401227_rf
 TGGGCAATTACTATTCTGATGAGTTCTAGTGTGCACATTCACCAATGTGGATAATAAATAGG
 TTAACCTCTAGGATTTTGAGAATTGAAATAATATATATCTTCTAGCCAGTAAATCATTTAAATT
 TTTTTTTACTAAGCATTAGCTGAAATCATTGSTATTTGTTTATTCTCTATCTACCCCTAAACATTGA
 AGTGTCTGGACCGAAAGTTGATATTACATCTTCAAGAACCTCTACCATGGGGCCCTTTCCTT
 TATTAAATTCTGCTATAATTAAATTGCTCTATTATGCCATTTCATGTTCAAGACACACTCTAAA
 TGTATA
 AGATGCCAAACACCTGGTAAATGCAATTCCAAAGCTCAACATGGCTGAAATTCACTACTCTT
 GGGCTCAGTTTGCCTATCTGAAATGAGTGAATAGGAAAGTGTTATTAAATTAAATTAA
 TACTTTTATTTAGATTTGGGATAACATGTTAAGTTGCTATAAGGAAATTCAAGTCATGGGAGT
 TTAATGAAAGATATTTCATCCAGGTAACAAAGCTAAACCTAAAGTAAATTGTTTAA
 TTTTCCCTCCCAACTT
 TGTGTCTCATCATTTAGCCCCACCTTAAGTGATAATATGTTGT
 TGTGTCTCATCATTTAGCCCCACCTTAAGTGATAATATGTTGT

CLAIMS

1. A process for detecting a chromosome state which represents a disease subgroup in a population comprising determining whether a chromosome interaction relating to that chromosome state is present or absent within a defined region of the genome, wherein said disease subgroup is an amyotrophic lateral sclerosis (ALS) subgroup; and
 - wherein said chromosome interaction has optionally been identified by a method of determining which chromosomal interactions are relevant to a chromosome state corresponding to an ALS subgroup of the population, comprising contacting a first set of nucleic acids from subgroups with different states of the 10 chromosome with a second set of index nucleic acids, and allowing complementary sequences to hybridise, wherein the nucleic acids in the first and second sets of nucleic acids represent a ligated product comprising sequences from both the chromosome regions that have come together in chromosomal interactions, and wherein the pattern of hybridisation between the first and second set of nucleic acids allows a determination of which chromosomal interactions are specific to an ALS subgroup; and
 - 15 - wherein the chromosome interaction:
 - (i) is present in any one of the regions or genes listed in Table 1 or 5; and/or
 - (ii) corresponds to any one of the chromosome interactions represented by any probe shown in Table 1 or 5, and/or
 - (iii) corresponds to any one of the chromosome interactions shown in Table 10 or 11, and/or
 - 20 (iv) is present in a 4,000 base region which comprises or which flanks (i), (ii) or (iii).
2. A process according to claim 1 which is carried out to diagnose ALS or to determine a prognosis for ALS.
3. A process according to claim 1 or 2 wherein a specific combination of chromosome interactions are 25 typed:
 - (i) comprising all of the chromosome interactions represented by the probes in Table 1 or 5; or
 - (ii) comprising at least 4, 5, 6 or 7 of the chromosome interactions represented by the probes in Table 1 or 5; or
 - (iii) which together are present in at least 4, 5, 6 or 7 of the regions or genes listed in Table 1 or 5; or
 - 30 (iv) comprising at least 4, 5, 6 or 7 chromosome interactions which are present in a 4,000 base region which comprises or which flanks the chromosome interactions represented by the probes in Table 1 or 5; or
 - (v) comprising all of the chromosome interactions shown Table 10 or 11; or
 - (vi) comprising at least 4, 5, 6 or 7 of the chromosome interactions shown in Table 10.

4. A process according to any one of the preceding claims in which the chromosome interactions are typed:

- in a sample from an individual, and/or
- by detecting the presence or absence of a DNA loop at the site of the chromosome interactions, and/or
- 5 - detecting the presence or absence of distal regions of a chromosome being brought together in a chromosome conformation, and/or
- by detecting the presence of a ligated nucleic acid which is generated during said typing and whose sequence comprises two regions each corresponding to the regions of the chromosome which come together in the chromosome interaction, wherein detection of the ligated nucleic acid is preferably by
- 10 using (i) a probe that has at least 70% identity to any of the specific probe sequences mentioned in Table 1 or 5, and/or (ii) by a primer pair which has at least 70% identity to any primer pair in Table 2 or 7.

5. A process according to any one of the preceding claims, wherein:

- 15 - the second set of nucleic acids is from a larger group of individuals than the first set of nucleic acids; and/or
- the first set of nucleic acids is from at least 8 individuals; and/or
- the first set of nucleic acids is from at least 4 individuals from a first subgroup and at least 4 individuals from a second subgroup which is preferably non-overlapping with the first subgroup; and/or
- 20 - the process is carried out to select an individual for a medical treatment.

6. A process according to any one of the preceding claims wherein:

- the second set of nucleic acids represents an unselected group; and/or
- wherein the second set of nucleic acids is bound to an array at defined locations; and/or
- 25 - wherein the second set of nucleic acids represents chromosome interactions in at least 100 different genes; and/or
- wherein the second set of nucleic acids comprises at least 1,000 different nucleic acids representing at least 1,000 different chromosome interactions; and/or
- wherein the first set of nucleic acids and the second set of nucleic acids comprise at least 100 nucleic acids with length 10 to 100 nucleotide bases.
- 30

7. A process according to any one of the preceding claims, wherein the first set of nucleic acids is obtainable in a process comprising the steps of: -

- (i) cross-linking of chromosome regions which have come together in a chromosome interaction;

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- (ii) subjecting said cross-linked regions to cleavage, optionally by restriction digestion cleavage with an enzyme; and
- (iii) ligating said cross-linked cleaved DNA ends to form the first set of nucleic acids (in particular comprising ligated DNA).

5 8. A process according to any one of the preceding claims wherein at least 5 to 9 different chromosome interactions are typed, preferably in 5 to 9 different regions or genes.

9. A process according to any one of the preceding claims wherein said defined region of the genome:

(i) comprises a single nucleotide polymorphism (SNP); and/or

(ii) expresses a microRNA (miRNA); and/or

10 (iii) expresses a non-coding RNA (ncRNA); and/or

(iv) expresses a nucleic acid sequence encoding at least 10 contiguous amino acid residues; and/or

(v) expresses a regulating element; and/or

(vii) comprises a CTCF binding site.

15 10. Method for identifying or designing a therapeutic agent for treating ALS by selecting an agent that is able to cause a change of chromosomal interaction and thereby cause a therapeutic effect,

- wherein the chromosomal interaction is represented by any probe in Table 1 or 5; and/or

- the chromosomal interaction is present in any region or gene listed in Table 1 or 5; and/or

20 - the chromosomal interaction is any one of the interactions shown in Table 10 or 11,

and wherein optionally:

- the chromosomal interaction has been identified by the method of determining which chromosomal interactions are relevant to a chromosome state as defined in claim 1, and/or
- the change in chromosomal interaction is monitored using (i) a probe that has at least 70% identity to any of the probe sequences mentioned in Table 1 or 5, and/or (ii) by a primer pair which has at least 70% identity to any primer pair in Table 1 or 5; and/or
- a candidate agent is contacted with a cell and the chromosome interaction in the cell is monitored to determine whether the candidate agent is able to treat ALS.

25 30 11. Use of

(iv) detection of a chromosomal interaction wherein:

- the chromosomal interaction is as represented by a probe in Table 1 or 5, and/or

- the chromosomal interaction is present in any region or gene which is mentioned in Table 1 or 5; and/or

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- the chromosomal interaction is any one of the interactions shown in Table 10 or 11;
- or
- (v) a probe that has at least 70% identity to any of the probe sequences mentioned in Table 1 or 5, or
- 5 (vi) a primer pair which has at least 70% identity to any primer pair identified in Table 2 or 7; to identify or design a therapeutic agent for ALS.
- 12. Use according to claim 11 to identify a therapeutic agent comprising administering a candidate agent, and using said detection of a chromosomal interaction, said probe or said primer pair to detect whether there is a change in chromosome state to thereby determine whether the candidate agent is a therapeutic agent, wherein the use is optionally performed *in vitro*, preferably in a cell.
- 13. A therapeutic agent for ALS for use in a method of preventing or treating ALS in an individual that has been identified as being in need of the therapeutic agent by a process according to any one of claims 1 to 9.
- 14. A process, method or use according to any of the preceding claims wherein the typing or detecting comprises specific detection of the ligated product by quantitative PCR (qPCR) which uses primers capable of amplifying the ligated product and a probe which binds the ligation site during the PCR reaction, wherein said probe comprises sequence which is complementary to sequence from each of the chromosome regions that have come together in the chromosome interaction, wherein preferably said probe comprises:
 - an oligonucleotide which specifically binds to said ligated product, and/or
 - a fluorophore covalently attached to the 5' end of the oligonucleotide, and/or
 - a quencher covalently attached to the 3' end of the oligonucleotide, and
- 25 optionally
 - said fluorophore is selected from HEX, Texas Red and FAM; and/or
 - said probe comprises a nucleic acid sequence of length 10 to 40 nucleotide bases, preferably a length of 20 to 30 nucleotide bases.
- 30 15. A process for detecting a chromosome state which represents a disease subgroup in a population comprising determining whether a chromosome interaction relating to that chromosome state is present or absent within a defined region of the genome, wherein said disease subgroup is a Huntington's disease subgroup; and
 - wherein said chromosome interaction has optionally been identified by a method of determining which chromosomal interactions are relevant to a chromosome state corresponding to an Huntington's disease

subgroup of the population, comprising contacting a first set of nucleic acids from subgroups with different states of the chromosome with a second set of index nucleic acids, and allowing complementary sequences to hybridise, wherein the nucleic acids in the first and second sets of nucleic acids represent a ligated product comprising sequences from both the chromosome regions that have come together in 5 chromosomal interactions, and wherein the pattern of hybridisation between the first and second set of nucleic acids allows a determination of which chromosomal interactions are specific to an Huntington's disease subgroup; and

- wherein the chromosome interaction:

(i) is present in any one of the regions or genes listed in Table 12; and/or

10 (ii) corresponds to any one of the chromosome interactions represented by any probe shown in Table 12, and/or

(iii) corresponds to any one of the chromosome interactions represented in Table 12, and/or

(iii) is present in a 4,000 base region which comprises or which flanks (i), (ii) or (iii).

15 16. A process according to claim 15 which is carried out to diagnose Huntington's disease or to determine a prognosis for Huntington's disease.

17. A process according to claim 15 or 16 wherein a specific combination of chromosome interactions are typed:

20 (i) comprising all of the chromosome interactions represented by the probes in Table 12; or

(ii) comprising at least 4, 5, 6 or 7 of the chromosome interactions represented by the probes in Table 12; or

(iii) which together are present in at least 4, 5, 6 or 7 of the regions or genes listed in Table 12; or

(iv) at least 4, 5, 6 or 7 chromosome interactions are typed which are present in a 4,000 base region which

25 comprises or which flanks the chromosome interactions represented by the probes in Table 12.

18. A process according to any one of claims 15 to 17 in which the chromosome interactions are typed:

- in a sample from an individual, and/or

- by detecting the presence or absence of a DNA loop at the site of the chromosome interactions, and/or

30 - detecting the presence or absence of distal regions of a chromosome being brought together in a chromosome conformation, and/or

- by detecting the presence of a ligated nucleic acid which is generated during said typing and whose sequence comprises two regions each corresponding to the regions of the chromosome which come together in the chromosome interaction, wherein detection of the ligated nucleic acid is preferably by

35 using (i) a probe that has at least 70% identity to any of the specific probe sequences mentioned in Table 12, and/or (ii) by a primer pair which has at least 70% identity to any primer pair in Table 12.

19. A process according to any one of claims 15 to 18, wherein:

- the second set of nucleic acids is from a larger group of individuals than the first set of nucleic acids; and/or

5 - the first set of nucleic acids is from at least 8 individuals; and/or

- the first set of nucleic acids is from at least 4 individuals from a first subgroup and at least 4 individuals from a second subgroup which is preferably non-overlapping with the first subgroup; and/or

- the process is carried out to select an individual for a medical treatment.

10 20. A process according to any one of claims 15 to 19 wherein:

- the second set of nucleic acids represents an unselected group; and/or

- wherein the second set of nucleic acids is bound to an array at defined locations; and/or

- wherein the second set of nucleic acids represents chromosome interactions in at least 100 different genes; and/or

15 - wherein the second set of nucleic acids comprises at least 1,000 different nucleic acids representing at least 1,000 different chromosome interactions; and/or

- wherein the first set of nucleic acids and the second set of nucleic acids comprise at least 100 nucleic acids with length 10 to 100 nucleotide bases.

20 21. A process according to any one of the claims 15 to 20, wherein the first set of nucleic acids is obtainable in a process comprising the steps of: -

(i) cross-linking of chromosome regions which have come together in a chromosome interaction;

(ii) subjecting said cross-linked regions to cleavage, optionally by restriction digestion cleavage with an enzyme; and

25 (iii) ligating said cross-linked cleaved DNA ends to form the first set of nucleic acids (in particular comprising ligated DNA).

22. A process according to any one of claims 15 to 21 wherein at least 5 to 9 different chromosome interactions are typed.

23. A process according to any one of claims 15 to 22 wherein said defined region of the genome:

30 (i) comprises a single nucleotide polymorphism (SNP); and/or

(ii) expresses a microRNA (miRNA); and/or

(iii) expresses a non-coding RNA (ncRNA); and/or

(iv) expresses a nucleic acid sequence encoding at least 10 contiguous amino acid residues; and/or

(v) expresses a regulating element; and/or

(vii) comprises a CTCF binding site.

24. Method for identifying or designing a therapeutic agent for treating Huntington's disease by selecting an agent that is able to cause a change of chromosomal interaction and thereby cause a therapeutic effect,

- wherein the chromosomal interaction is represented by any probe in Table 12; and wherein optionally:
 - the chromosomal interaction has been identified by the method of determining which chromosomal interactions are relevant to a chromosome state as defined in claim 15, and/or
 - the change in chromosomal interaction is monitored using (i) a probe that has at least 70% identity to any of the probe sequences mentioned in Table 12, and/or (ii) by a primer pair which has at least 70% identity to any primer pair in Table 12; and/or
 - a candidate agent is contacted with a cell and the chromosome interaction in the cell is monitored to determine whether the candidate agent is able to treat Huntington's disease.

15 25. Use of

- detection of a chromosomal interaction wherein the chromosomal interaction is as represented by a probe in Table 12; or
 - a probe that has at least 70% identity to any of the probe sequences mentioned in Table 12, or
 - a primer pair which has at least 70% identity to any primer pair identified in Table 12;

20 to identify or design a therapeutic agent for Huntington's disease.

26. Use according to claim 25 to identify a therapeutic agent comprising administering a candidate agent, and using said detection of a chromosomal interaction, said probe or said primer pair to detect whether there is a change in chromosome state to thereby determine whether the candidate agent is a therapeutic agent, wherein the use is optionally performed *in vitro*, preferably in a cell.

25

27. A therapeutic agent for Huntington's disease for use in a method of preventing or treating Huntington's disease in an individual that has been identified as being in need of the therapeutic agent by a process according to any one of claims 15 to 23.

30 28. A process, method or use according to any of the preceding claims wherein the typing or detecting comprises specific detection of the ligated product by quantitative PCR (qPCR) which uses primers capable of amplifying the ligated product and a probe which binds the ligation site during the PCR reaction, wherein said probe comprises sequence which is complementary to sequence from each of the chromosome regions that have come together in the chromosome interaction, wherein preferably said probe comprises:

an oligonucleotide which specifically binds to said ligated product, and/or
a fluorophore covalently attached to the 5' end of the oligonucleotide, and/or
a quencher covalently attached to the 3' end of the oligonucleotide, and
optionally

5 said fluorophore is selected from HEX, Texas Red and FAM; and/or
said probe comprises a nucleic acid sequence of length 10 to 40 nucleotide bases, preferably a length of
20 to 30 nucleotide bases.

10

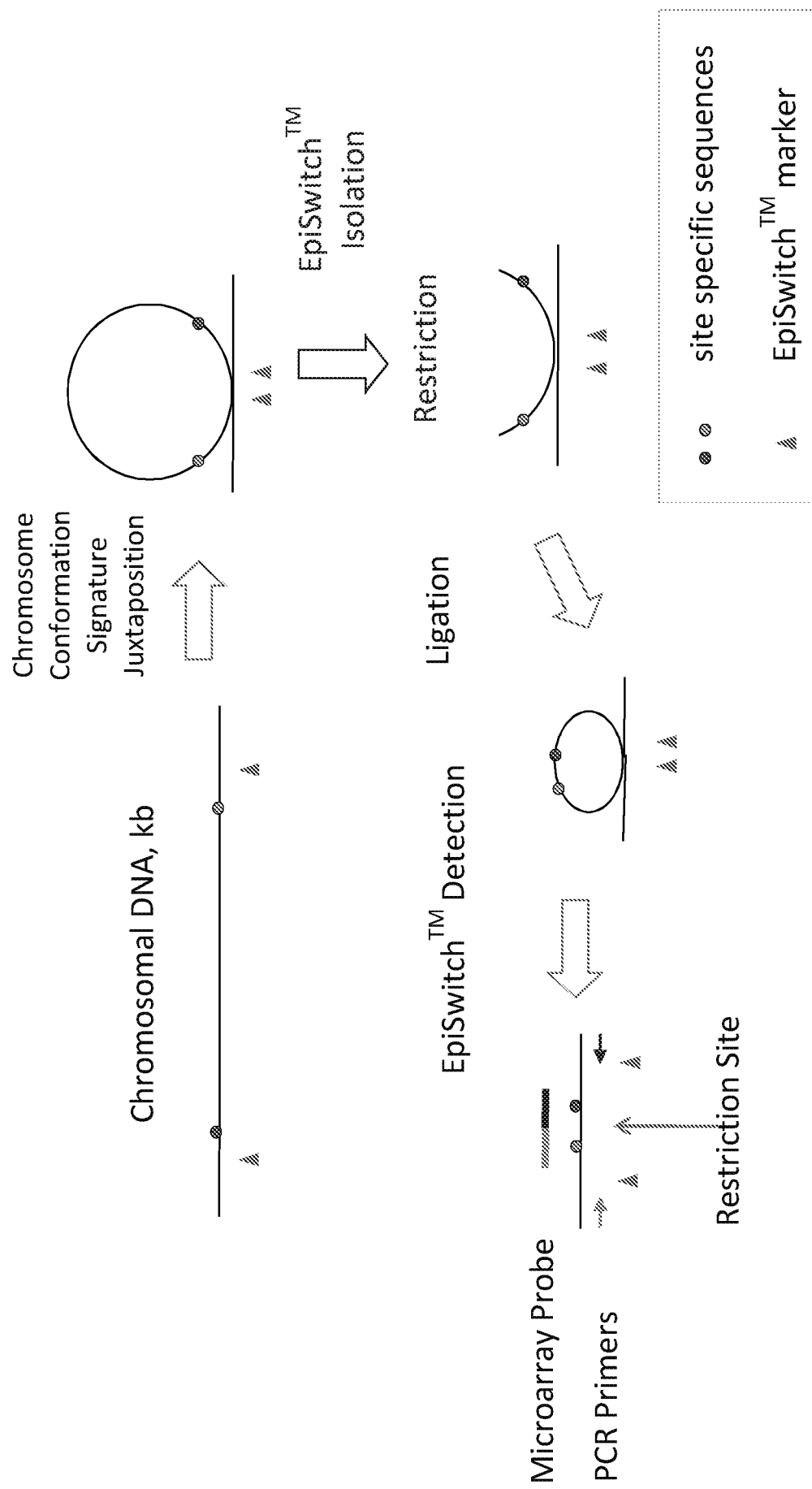
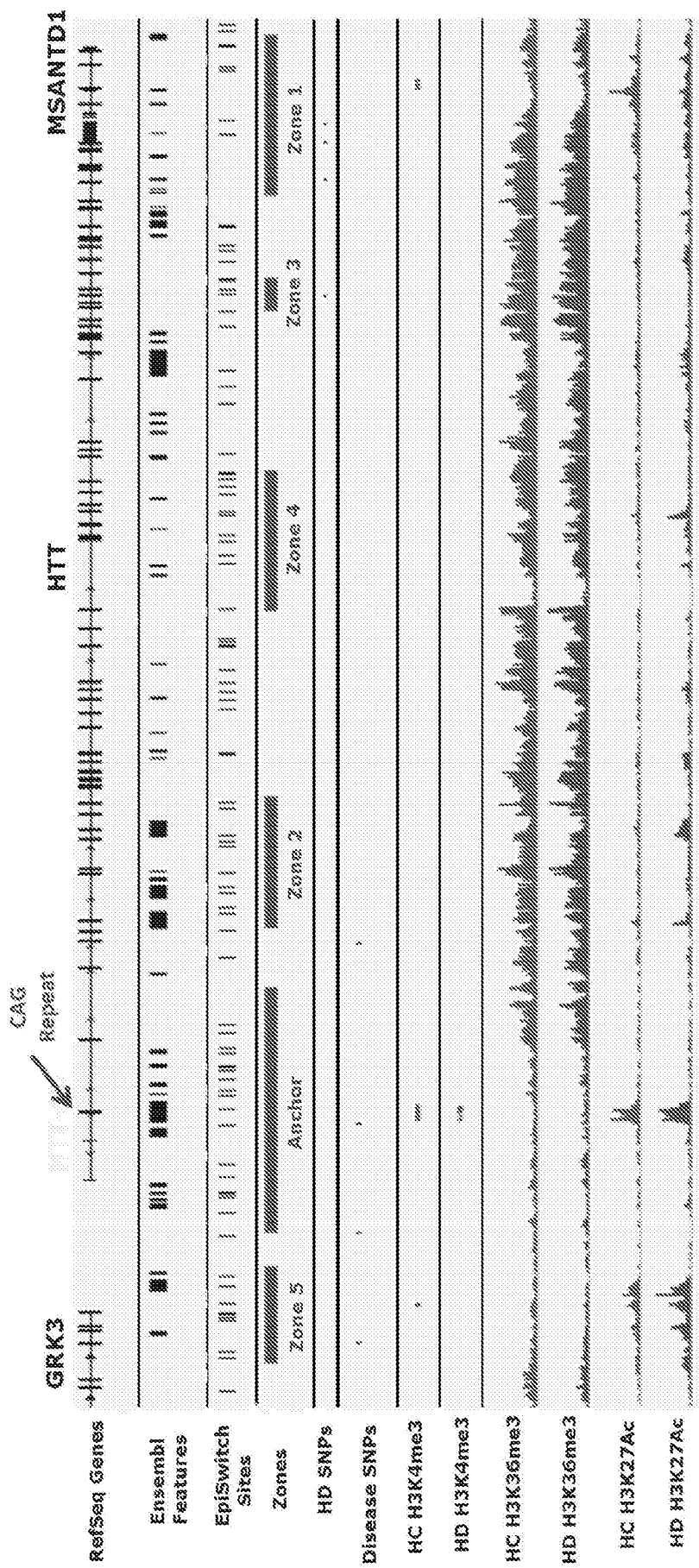
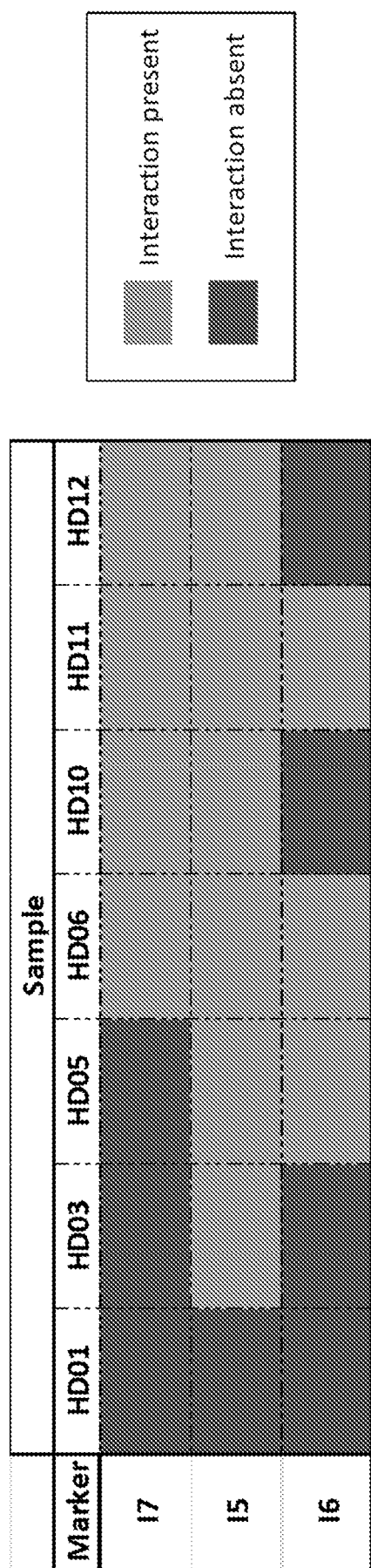


Figure 1.

Figure 2.



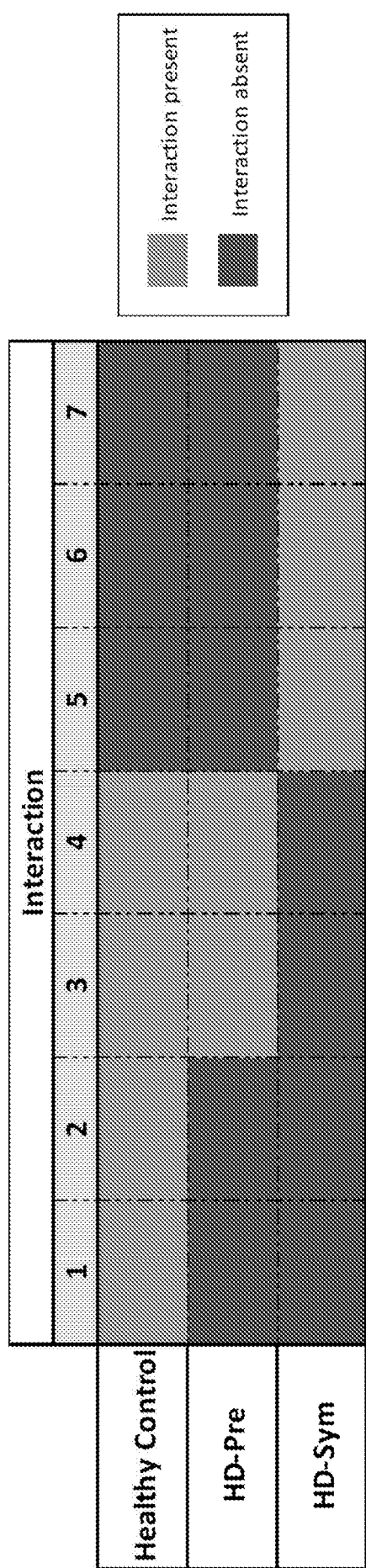


Figure 4.

Figure 5.