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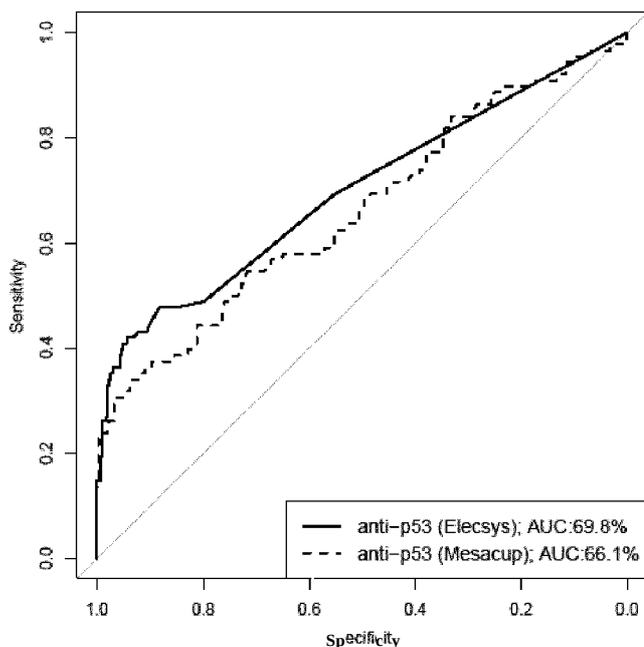
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(54) Title: DETECTION OF ANTI-P53 ANTIBODIES

Figure 2



(57) Abstract: The disclosure relates to an in vitro method for detecting an antibody to p53 (anti-p53 antibody) in a sample, the method comprising: incubating a sample to be analyzed with a p53 capture antigen and a p53 detection antigen, whereby a complex comprising the p53 capture antigen, the anti-p53 antibody and the p53 detection antigen is formed, separating the complex formed from unbound detection antigen and measuring the complex obtained via the detection antigen comprised therein, thereby detecting the anti-p53 antibody comprised in the sample.

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Detection of anti-p53 antibodies

Background of the Invention

The present disclosure relates to a method for the detection of antibodies against p53 (anti-p53 antibodies). The method disclosed, enables the detection of anti-p53 antibodies with higher specificity as compared to state of the art methods. The method according to the present disclosure requires a p53 capture antigen and a p53 detection antigen. The method comprises incubating a p53 capture antigen and a p53 detection antigen with a sample known or suspected to comprise an anti-p53 antibody, whereby a complex comprising the p53 capture antigen the anti-p53 antibody and the p53 detection antigen is formed, separating the complex formed from unbound detection antigen, measuring the complex obtained via the detection antigen comprised therein and thereby detecting the anti-p53 antibody comprised in the sample.

Anti-p53 antibodies (anti-p53 Abs) were discovered more than 30 years ago during the course of tumor-associated antigen screening, Crawford L.V. et al, *Int. J. Cancer*, 30: 403-408, 1982. However, this study as well as a few similar ones, performed in the early 1980s, were virtually ignored for more than 10 years because of a lack of interest in p53 during that period.

In the early 1990s, it was discovered that the p53 gene is the most common target for molecular alteration in almost every type of human cancer. This provoked considerable interest in the study of the p53 protein and its function in normal and transformed cells. It also led to the rediscovery of the humoral response, which had been found earlier already in cancer patients.

The tumor suppressor p53 is a phosphoprotein barely detectable in the nucleus of normal cells (Benchimol, S. et al, *EMBO J.* 1 (1982) 1055-1062). On cellular stress, particularly that induced by DNA damage, p53 can arrest cell cycle progression, thus allowing the DNA to be repaired or it can lead to apoptosis. In cancer cells that bear a mutant p53, this protein is no longer able to control cell proliferation, which results in inefficient DNA repair (Levine, A. *Cell* 88 (1997) 323-331). The most common changes of p53 in human cancer are point missense mutations, which are found in cancers of the colon, stomach, breast, lung, brain and esophagus (Greenblatt, M. et al, *Cancer Res.* 54 (1994) 4855-4878). It is estimated that p53 mutations is the most frequent genetic event in human cancers and

accounts for more than 50% of cases. There is a very strong correlation between the frequency of p53 antibodies and the frequency of p53 mutations arguing that p53 mutations are involved in the appearance of these antibodies (Soussi, T. Cancer Res. 60 (2000) 1777-1788). Anti-p53 antibodies are found in human cancer patients with a specificity of about 96%, but the sensitivity of such detection is only about 30%.

The association of p53 mutations with anti-p53 antibodies suggests that the anti-p53 humoral immune response is due to a self-immunization process linked to the strong immunogenicity of the p53 protein.

The role of p53 mutations in carcinogenesis has been extensively studied and anti-p53 antibodies have been found to be associated to numerous cancer types such as Colorectal Cancer (CRC), Oesophageal Cancer and Breast Cancer among others. These 3 cancer types represent a considerable healthcare burden.

With an incidence of 746,000 cases/year in men, CRC is the third most common cancer in men worldwide and represents 10.0% of the total cancers among male patients. CRC is also the second most common in women across the world with 614,000 new cases each year, representing 9.2% of cancers affecting female patients. Oesophageal cancer is the sixth most common cause of death from cancer and has an estimated incidence of 456,000 new cases among men and women representing 3.2% of the total new cancers each year. Breast Cancer is by far the most frequent cancer among women with an estimated 1.67 million new cancer cases diagnosed worldwide in 2012 representing 25% of all cancers in female patients. Overall, Breast cancer is the second most common cancer in the world. As obvious from the high incidence of cancer types associated with anti-p53 (auto-) antibodies, there is a need for a sensitive and reliable diagnostic method for detection of anti-p53 (auto-)antibodies.

As summarized by Soussi, T. (Cancer Research 60 (2000) 1777-1788) many papers on anti-p53 have been published. The detection methods used vary significantly with ELISA-methods, immuno precipitation methods and Western Blotting being the most prominent ones. It may well be that the use of many different methods for detection of anti-p53 antibodies has lead to some discrepant results and to some uncertainty regarding the diagnostic utility of anti-p53 antibodies.

The availability of the first commercial assays for measurement of anti-p53 antibodies has been of great value in improving the comparability of data/papers on

anti-p53 antibodies. One of the most frequently used assay for measurement of anti-p53 antibodies is "MESACUP Anti-p53 TEST" distributed by MEDICAL & BIOLOGICAL LABORATORIES CO., LTD., KDX Nagoya, Japan. This Enzyme immunoassay for the detection of anti-p53 IgG in human serum is laborious and
5 e.g. requires the parallel measurement of both specific (including non-specific) and non-specific binding, respectively. The fraction of specific binding has then to be calculated by subtraction of the value determined for non-specific binding from the sum of specific and non-specific binding.

Antibodies of the IgM class of antibodies in general are more sticky and lead to
10 further complications, e.g. further cross-reactivity and non-specific binding. For this reason anti-p53 detection methods in most cases rely on anti-human IgG detection reagents, i.e. the detection of only the human (auto-) antibodies of the immunoglobulin G class and not those of the immunoglobulin M class. As further evidenced by the way the Mesacup assay has to be performed, non-specific binding
15 most likely is a key problem for any method used in the detection of anti-p53 antibodies.

As obvious from the high incidence of cancer types associated with anti-p53 (auto-) antibodies, and as further evident from the drawbacks with state of the art methods used to detect those antibodies there is a tremendous need for a sensitive and
20 reliable immuno assay method for detection of anti-p53 (auto-)antibodies.

It surprisingly has been found that the method for detecting anti-p53 (auto-) antibodies as disclosed herein below is rapid, reliable, specific, sensitive and has the potential to set a new standard in the detection of anti-p53 (auto-) antibodies.

Summary of the Invention

25 Herein is reported an in vitro method for detecting an antibody to p53 (anti-p53 antibody) in a sample, the method comprising: a) incubating a sample to be analyzed with a p53 capture antigen and a p53 detection antigen, whereby a complex comprising the p53 capture antigen, the anti-p53 antibody and the p53
30 detection antigen is formed, b) separating the complex formed in (a) from unbound detection antigen, c) measuring the complex obtained in step (b) via the detection antigen comprised therein, thereby detecting the anti-p53 antibody comprised in the sample. Further and more detailed embodiments relate, e.g., to specific partial sequences of p53 for use in a method according to the present disclosure.

Detailed Description of the Invention

The present disclosure relates to an in vitro method for detecting an antibody to p53 (anti-p53 antibody) in a sample, the method comprising: a) incubating a sample to be analyzed with a p53 capture antigen and a p53 detection antigen, whereby a complex comprising the p53 capture antigen, the anti-p53 antibody and the p53 detection antigen is formed, b) separating the complex formed in (a) from unbound detection antigen, c) measuring the complex obtained in step (b) via the detection antigen comprised therein, thereby detecting the anti-p53 antibody comprised in the sample.

10 In the following the terms "anti-p53 (auto-) antibody", "anti-p53 antibody" or "p53 antibody" are used interchangeably. An anti-p53 antibody is an antibody that binds to the p53 polypeptide of SEQ ID NO: 3 or to a partial sequence thereof.

As mentioned above the method according to the present invention requires the use of both, a p53 capture antigen and of a p53 detection antigen, respectively. An antibody to p53 present in a sample to be investigated binds to both these antigens, thereby forming a complex comprising the p53 capture antigen, the anti-p53 antibody and the p53 detection antigen.

The term "capture antigen" is familiar to a person skilled in the art.

In order to perform a double antigen sandwich immuno assay it is required that one provides an epitope at least twice, at least once on the capture antigen to mediate binding to a solid phase, and at least once on the detection antigen to allow for detection of the sandwich complex (antigen-antibody-antigen) formed.

In one embodiment the p53 capture antigen is capable of binding to a solid phase.

Materials for manufacturing a solid phase (solid support) are well-known in the art. and include, inter alia, commercially available column materials, polystyrene beads, latex beads, magnetic beads, colloid metal particles, glass and/or silicon chips and surfaces, nitrocellulose strips, membranes, sheets, duracytes, wells and walls of reaction trays, wells and walls of reaction vessels, plastic tubes etc.

30 In one embodiment the solid phase is represented by the wells and walls of a microtitre plate.

In one embodiment the solid phase is in the form of particles. The particle may be a microparticle which is or which comprises magnetic or ferromagnetic metals, alloys or compositions. In further embodiments, the solid phase material may have specific properties and e.g. be hydrophobic, or hydrophilic. In one embodiment of the present invention, the microparticles are paramagnetic microparticles and the separation of such particles in the measurement method according to the present disclosure is facilitated by magnetic forces. Magnetic forces are applied to pull the paramagnetic or magnetic particles out of the solution/suspension and to retain them as desired while liquid of the solution/suspension can be removed and the particles can e.g. be washed.

As indicated above, in one embodiment the p53 capture antigen is capable of binding to a solid phase. In order for the capture antigen to be capable of binding to a solid phase it is preferred to employ a specific binding pair. One partner of such binding pair is bound to the capture antigen and the other partner of such binding pair is bound to the solid phase.

A "binding pair" as used herein consists of two partners binding to each other with high affinity, i.e. with one nanomolar affinity or better. Embodiments for binding pairs are for example the binding pairs consisting of receptor and ligand, antigen and antibody, hapten and anti-hapten antibody, and binding pairs based on naturally occurring high affinity binding pairs.

One example of a receptor-ligand binding pair is a pair consisting of a steroid hormone receptor and the corresponding steroid hormone.

One further type of a binding pair which is suitable for the method according to the present invention is a hapten and anti-hapten antibody binding pair. A hapten is an organic molecule with a molecular weight of 100 to 2000 Dalton, preferably of 150 to 1000 Dalton. Such small molecule which if used as such is non-immunogenic but can be rendered immunogenic by coupling it to a carrier molecule and anti-hapten antibodies can be generated according to standard procedures. The hapten may be selected from the group comprising sterols, bile acids, sexual hormones, corticoids, cardenolides, cardenolide-glycosides, bufadienolides, steroid-sapogenines and steroid alkaloids, cardenolides and cardenolide-glycosides. Representatives of these substance classes are digoxigenin, digitoxigenin, gitoxigenin, strophanthidin, digoxin, digitoxin, ditoxin, strophanthin. Another suitable hapten is for example fluorescein.

5 Examples of binding pairs based on naturally occurring high affinity binding pairs are biotin or biotin analogues such as aminobiotin, iminobiotin or desthiobiotin and avidin or streptavidin as well as the FimG and DsF binding pair. The biotin-(strept)avidin binding pair is well-known in the art. The basic principles of the FimG-DsF binding pair are e.g. described in WO2012/028697.

In one embodiment the p53 capture antigen is covalently bound to a first partner of a binding pair and the second partner of said binding pair is bound to the solid phase.

10 In one embodiment the first partner of the binding pair is selected from hapten, biotin or a biotin analogue such as aminobiotin, iminobiotin or desthiobiotin, DsF and the ligand for a receptor and the second partner of said binding pair is selected from anti-hapten antibody, avidin or streptavidin, FimG and a ligand-receptor.

15 In one embodiment first partner of the binding pair is selected from hapten, biotin or a biotin analogue such as aminobiotin, iminobiotin or desthiobiotin and DsF, and the second partner of said binding pair is selected from anti-hapten antibody, avidin or streptavidin and FimG.

20 In one embodiment first partner of the binding pair is selected from biotin or a biotin analogue, such as aminobiotin, iminobiotin or desthiobiotin, and DsF, and the second partner of said binding pair is selected from, avidin or streptavidin and FimG.

In one embodiment first partner of the binding pair is biotin or a biotin analogue such as aminobiotin, iminobiotin or desthiobiotin and the second partner of said binding pair is selected from avidin or streptavidin.

25 In one embodiment first partner of the binding pair is biotin and the second partner of said binding pair is streptavidin.

In one embodiment the method according to the present invention is practiced with a p53 capture antigen that is biotinylated and a solid phase that is coated with avidin or streptavidin.

30 It is, however, also possible to directly bind the capture antigen to a solid phase. In one embodiment the method according to the present invention is practiced with a p53 capture antigen that is bound to the solid phase.

Antibodies are at least bi-valent, i.e. having the potential to bind with one 'arm' to the capture antigen and with the other arm (or any of the other arms for multivalent antibodies) to the detection antigen.

5 An antibody to p53 present in a sample to be investigated binds to both these antigens, thereby forming a complex comprising the capture antigen, the anti-p53 antibody and the detection antigen.

The skilled artisan is familiar with the term "detection antigen".

10 The term detection antigen implies that this antigen can be detected. Such detection can be accomplished by various types of labels which are either directly or indirectly detectable.

In one embodiment the method according to the present disclosure is a method wherein the p53 detection antigen comprises a directly detectable label.

The term detectably labeled encompasses labels that can be directly or indirectly detected.

15 Directly detectable labels either provide a detectable signal or they interact with a second label to modify the detectable signal provided by the first or second label, e.g. to give FRET (fluorescence resonance energy transfer). Labels such as fluorescent dyes and luminescent (including chemiluminescent and electrochemiluminescent) dyes (Briggs et al "Synthesis of Functionalised
20 Fluorescent Dyes and Their Coupling to Amines and Amino Acids," J. Chem. Soc, Perkin-Trans. 1 (1997) 1051-1058) provide a detectable signal and are generally applicable for labeling. In one embodiment detectably labeled refers to a label providing or inducible to provide a detectable signal, i.e. to a luminescent label, to a fluorescent label, to a chemiluminescent label or to an electrochemiluminescent
25 label, respectively.

In one embodiment according to the present disclosure the microparticle-based analyte-specific binding assay makes use of a chemiluminescent or an electrochemiluminescent label and a corresponding light detection system. The light produced by the label is measured and directly or indirectly indicates the
30 presence or quantity of the analyte.

Electrochemiluminescent (ECL) assays provide a sensitive and precise measurement of the presence and concentration of an analyte of interest. Such

techniques use labels or other reactants that can be induced to luminesce when electrochemically oxidized or reduced in an appropriate chemical environment. Such electrochemiluminescence is triggered by a voltage imposed on a working electrode at a particular time and in a particular manner. The light produced by the label is measured and indicates the presence or quantity of the analyte. For a fuller description of such ECL techniques, reference is made to US Patent No. 5,221,605, US Patent No. 5,591,581, US Patent No. 5,597,910, PCT published application WO90/05296, PCT published application W092/14139, PCT published application WO90/05301, PCT published application WO96/24690, PCT published application US95/03190, PCT application US97/16942, PCT published application US96/06763, PCT published application WO95/08644, PCT published application WO96/06946, PCT published application W096/3341 1, PCT published application WO87/06706, PCT published application W096/39534, PCT published application W096/41 175, PCT published application WO96/40978, PCT/US97/03653 and US patent application 08/437,348 (U.S. Patent No. 5,679,519). Reference is also made to a 1994 review of the analytical applications of ECL by Knight, et al. (Analyst, 1994, 119: 879-890) and the references cited therein. In one embodiment the method according to the present description is practiced using an electrochemiluminescent label.

In one embodiment the directly detectable label (used to label the detection antigen in a method of the present disclosure) is selected from the group consisting of a luminescent label, a fluorescent label, a chemiluminescent label or an electrochemiluminescent label.

In one embodiment the directly detectable label is a chemiluminescent or an electrochemiluminescent label.

In one embodiment the double-antigen sandwich assay according to the present disclosure uses the full length p53 polypeptide of SEQ ID NO:3 both as the capture as well as the detection antigen.

A synthetic peptide, representing at least one important epitope of p53 of the p53 polypeptide is of high utility in a method according to the present disclosure. Without wanting to be bound to theory, it may well be that such synthetic peptide sequence comprises an epitope that is bound by a significant percentage of those antibodies in a sample that also bind to the full-length p53 polypeptide. At the same time it may well be that those stretches of the p53 polypeptide that are rarely or not

at all bound by an anti-p53 antibody (as e.g. comprised in a bodily fluid sample) are those sequence stretches that significantly contribute to the non-specific binding of antibodies, especially of the sticky IgM-class antibodies.

5 In one embodiment the in vitro method for detection of anti-p53 antibodies according to the present invention is a double antigen sandwich immuno assay method wherein both the p53 capture antigen and the p53 detection antigen comprise a peptide of SEQ ID NO: 1.

10 In one embodiment the in vitro method for detection of anti-p53 antibodies according to the present invention is a double antigen sandwich immuno assay method wherein both the p53 capture antigen and the p53 detection antigen comprise a peptide of SEQ ID NO: 2.

15 In one embodiment the method for detection of anti-p53 antibodies according to the present invention is a double antigen sandwich immuno assay method wherein the peptides of SEQ ID NO:1 and of SEQ ID NO:2 are each comprised in both the p53 capture antigen and the p53 detection antigen.

In one embodiment the peptides representing the relevant partial sequences of the p53 polypeptide used in a method for detection of anti-p53 antibodies according to the present invention are the peptides consisting of SEQ ID NO:1 and of SEQ ID NO:2, respectively.

20 In one embodiment the method for detection of anti-p53 antibodies according to the present invention is a double antigen sandwich immuno assay method wherein the p53 detection antigen is a single polypeptide comprising both the peptides of SEQ ID NO: 1 and of SEQ ID NO:2 at least twice .

25 In one embodiment the method for detection of anti-p53 antibodies according to the present invention is a double antigen sandwich immuno assay method wherein the p53 detection antigen comprises a fusion polypeptide comprising at least one multimerization domain, at least one polypeptide of SEQ ID NO: 1 and at least one polypeptide of SEQ ID NO:2, wherein the multimerization domain is a prokaryotic or eukaryotic chaperone selected from the group consisting of FkpA, Skp, SecB, Hsp25, MIP, GroEL, ClpB and ClpX.

30

In one embodiment the present disclosure relates to a fusion polypeptide comprising at least one multimerization domain, at least one polypeptide of SEQ

ID NO:1 and at least one polypeptide of SEQ ID NO:2, wherein the multimerization domain is a prokaryotic or eukaryotic chaperone selected from the group consisting of FkpA, Skp, SecB, Hsp25, MIP, GroEL, ClpB and ClpX.

5 The multimerization domain is preferably located at the N- and/or C-terminus of the fusion polypeptide, more preferably at the N-terminus. The multimerization domain is a polypeptide sequence which supports multimerization of individual fusion polypeptide molecules, wherein a multimer is formed which is comprised of a plurality of monomeric subunits, which are associated by non-covalent interactions. The monomeric subunits of the complex are genetic fusion proteins,
10 wherein the individual amino acid residues are linked by peptide bonds. The monomeric subunits of the multimer are preferably identical.

For example, the multimerization domain may be a dimerization domain, i.e. a domain which supports noncovalent association of two subunits, a trimerization domain, which supports non-covalent association of three subunits, a
15 tetramerization domain or an even higher multimerization domain. Preferably, the multimerization domain is a dimerization domain, a trimerization domain or a tetramerization domain.

Multimerization domains may be selected from prokaryotic or eukaryotic chaperones, preferably from ATPindependent chaperones. Specific examples of
20 multimerization domains are the proteins FkpA, Skp and SecB from *E. coli* or orthologs thereof from other prokaryotic organisms. FkpA is an ATP-independent periplasmic dimerization chaperone from *E. coli*. Skp is an ATP-independent periplasmic trimerization chaperone from *E. coli*. SecB is an ATPindependent cytosolic tetramerization chaperone from *E. coli*. Further suitable multimerization
25 domains are heat shock proteins from eukaryotic or prokaryotic organisms, e.g. Hsp25, an ATP-independent eukaryotic cytosolic/nuclear oligomeric chaperone. A further suitable multimerization domain is MIP (macrophage infectivity potentiator), an ATP-independent dimerization chaperone which is structurally related with FkpA. ATP-dependent chaperones like GroEL, an ATP-dependent
30 cytosolic heptamerisation chaperone from *E. coli* or ClpB, an ATP-dependent hexamerization chaperone from *E. coli* or ClpX are also suitable. Further, the multimerization domains may be selected from fragments or variants of the above polypeptides, which retain their ability of multimer formation.

The individual polypeptide sequences of SEQ ID NO:1 and of SEQ ID NO:2, respectively, in the fusion polypeptide may be separated by spacer sequences. The spacer sequences are preferably sequences which are heterologous to the p53 . For practical purposes, the spacer sequences are selected from sequences which barely
5 if at all interfere with the use of the fusion polypeptide as an antigen for determining anti-p53 antibodies. This means that the spacer sequences are not immunologically reactive against the antibodies to be tested. E.g., the spacer sequences comprise glycine and/or serine residues. In one embodiment the spacer is a poly-glycine spacer. The length of the spacer sequence is preferably from 1-10
10 amino acids.

Further, a spacer sequence may be present between the multimerization domain and the polypeptide of SEQ ID NO:1 and/or the polypeptide of SEQ ID NO:2.

This spacer sequence may have a length of e.g. 1-100 amino acids. Preferably, this spacer sequence is as described above.

15 In one embodiment the method according to the present disclosure is practiced with a p53 detection antigen comprising an indirectly detectable label.

Indirectly detectably labeled refers, e.g. to a detection antigen comprising a hapten and to the detection of such haptenylated compound by an anti-hapten antibody carrying a directly detectable label or to the a detection antigen comprising an
20 enzyme and to the detection of such enzyme by its corresponding enzymatic activity resulting in the conversion of an appropriate dye substrate. Various enzyme-substrate labels are available or disclosed (see e.g. US 4,275,149). The enzyme generally catalyzes a chemical alteration of a chromogenic substrate that can be measured using various techniques. For example, the enzyme may catalyze
25 a color change in a substrate, which can be measured spectrophotometrically. Alternatively, the enzyme may alter the fluorescence or chemiluminescence of the substrate. The chemiluminescent substrate becomes electronically excited by a chemical reaction and may then emit light which can be measured (using a chemiluminometer, for example) or donates energy to a fluorescent acceptor.
30 Examples of enzymatic labels include luciferases (e.g., firefly luciferase and bacterial luciferase; US 4,737,456), luciferin, 2,3-dihydrophthalazinediones, malate dehydrogenase, urease, peroxidase such as horseradish peroxidase (HRP), alkaline phosphatase (AP), (3-galactosidase, glucoamylase, lysozyme, saccharide oxidases (e.g., glucose oxidase, galactose oxidase, and glucose-6-phosphate dehydrogenase),

heterocyclic oxidases (such as uricase and xanthine oxidase), lactoperoxidase, microperoxidase, and the like. Techniques for conjugating enzymes to polypeptides are described in O'Sullivan et al "Methods for the Preparation of Enzyme-Antibody Conjugates for use in Enzyme Immunoassay", in *Methods in Enzym.* (ed. by J. Langone & IT Van Vunakis), Academic Press, New York, 73 (1981) 147-166.

Examples of enzyme-substrate combinations (US 4,275,149; US 4,318,980) include, for example: Horseradish peroxidase (HRP) with hydrogen peroxidase as a substrate, wherein the hydrogen peroxidase oxidizes a dye precursor (e.g., orthophenylene diamine (OPD) or 3,3',5,5'-tetramethylbenzidine hydrochloride (TMB)); alkaline phosphatase (AP) with para-nitrophenyl phosphate as chromogenic substrate; and (3-D-galactosidase ((3-D-Gal) with a chromogenic substrate (e.g., p-nitro phenyl-(3-D-galactosidase) or fluorogenic substrate 4-methylumbelliferyl-(3-D-galactosidase.

As mentioned above and as demonstrated in the Examples section, anti.p53 antibodies are most frequently found in CRC. In one embodiment the present invention relates to an in vitro method for diagnosing CRC by detecting an antibody to p53 (anti-p53 antibody) in a sample, the method comprising: a) incubating a sample to be analyzed with a p53 capture antigen and a p53 detection antigen, whereby a complex comprising the p53 capture antigen the anti-p53 antibody and the p53 detection antigen is formed, b) separating the complex formed in (a) from unbound detection antigen, c) measuring the complex obtained in step (b) via the detection antigen comprised therein, and d) diagnosing CRC if anti-p53 antibody is measured in the sample.

Liquid samples can be used in a method for specific in vitro-detection of anti-p53 antibodies in a method according to the present disclosure. The sample may be known to comprise an anti-p53 antibody or it may be suspected of comprising an anti-p53 antibody. In one embodiment the sample for in vitro diagnosis used in a method according to the present disclosure is a body fluid selected from whole blood, blood serum, blood plasma, liquor, urine or saliva. In one embodiment the sample suspected of comprising or comprising an anti-p53 antibody is serum, plasma or liquor. In one embodiment the sample suspected of comprising or comprising an anti-p53 antibody is serum or plasma.

The following examples and figures are provided to aid the understanding of the present invention, the true scope of which is set forth in the appended claims. It is

understood that modifications can be made in the procedures set forth without departing from the spirit of the invention.

Description of the Figures

Figure 1 Assay principle of Elecsys anti-p53 assay
5 The double antigen sandwich assay (DAGS) format is schematically depicted.
=> detection of anti-p53 auto-antibodies present in the sample is accomplished via biotinylated capture antigen (binding to streptavidin coated to paramagnetic beads) and ruthenylated p53
10 detection antigen.

Figure 2 ROC-Blots
For both assays analyzed the receiver-operator-curves (ROCs) are given. The ROC for the MesaCup assay is depicted as an interrupted line, whereas the ROC for the assays disclosed in the
15 present application is given as continuous/full line.

Example 1:

Capture and detection antigen used for detection of anti-p53 auto-antibodies

Various types of double antigen sandwich assay (DAGS) have been used.

20 It is possible to perform the DAGS assay for detection of anti-p53 auto-antibodies present in a sample as an enzyme-linked immuno sorbent assay (ELISA) in microtitre plates.

Synthetic peptides of SEQ ID NO:1 and SEQ ID NO:2, respectively proved to be advantageous. They could easily and reproducibly be synthesized and seemed to be appropriate due to the specific binding of anti-p53 antibodies thereto.

25 For higher through-put and higher reproducibility an anti-p53 assay was developed running on the automated electrochemiluminescence-based analyzers from Roche Diagnostics, Mannheim, Germany.

30 In one assays format two biotinylated peptides, one comprising the peptide of SEQ ID NO:1 and the other comprising the peptide of SEQ ID NO:2 are used as the capture antigen and the ruthenylated forms of both these peptides are used as the detection antigen.

Ruthenylation was performed according to standard procedures. In a typical protocol for ruthenylation the lysine ϵ -amino groups of the fusion protein are modified at protein concentrations of about 10 mg/ml with N-hydroxy-succinimide activated ruthenium labels. The label:protein molar ratio can be varied from 2:1 to 7:1, depending on the respective fusion protein an requirements for density of labels. The reaction buffer usually is 150 mM potassium phosphate (pH 8.0), 50 mM potassium chloride, 1 mM EDTA. The coupling or labeling reaction usually is carried out at room temperature for 10 minutes and stopped by adding buffered L-lysine to a final concentration of 10 mM. To avoid hydrolytic inactivation of the activated labels, the respective stock solutions are e.g. prepared in dried DMSO (seccosolv quality, Merck, Germany). DMSO concentrations up to 20 % in the reaction buffer are well tolerated by the fusion protein of SEQ ID NO:4 as disclosed herein. After the coupling reaction, unreacted free label and the organic solvent was removed by passing the crude protein conjugate over a gel filtration column (Superdex 200 HiLoad).

Due to the chaperon-part ("Skp") of the p53-construct, the protein spontaneously trimerizes during elution from the Ni-NTA column in the controlled refolding and purification process. The resulting trimer is very stable and remains its configuration during rhutenylation and further processing.

Example 2:

Elecsys® assay for measurement of anti-p53 auto-antibodies

The assay principle of the Elecsys® assay for detection of anti-p53 antibodies is depicted in Figure 1.

In brief, the biotinylated capture antigen (the biotinylated peptides of Eample 1.1), the sample and the ruthenylated detection antigen (the labeled fusion protein of Example 1.2) are incubated to allow for formation of a complex of capture antigen - anti-p53 antibody - detection antigen. Signal detection in the Elecsys® immunoassay is based on electrochemoluminescence. The biotin-conjugated capture antigen is immobilized on the surface of streptavidin-coated magnetic beads, whereas the detection antigen bears a complexed ruthenium cation as the luminescent moiety. In the presence of anti-p53 antibodies, the chromogenic ruthenium complex is bridged to the solid phase and emits light at 620 nm after excitation at a platinum electrode. The signal output is in arbitrary light units.

The following immunological reagents were used:

- 16 -

| Biotinylated capture antigen | Cone. | Ruthenylated detection antigen | Cone. |
|-------------------------------------|--------------|---------------------------------------|--------------|
| p53(1-35)[Bi(EU)3-1]amid | 20 ng/ml | rec.E.coli material | 300ng/ml |
| p53(41-60)[Bi(EU)3-41]amid | 20 ng/ml | Skp-(p53 peptides)x2-BPRU | |

75 μ l of the capture antigen comprising both peptides, each at the above given concentration, 75 μ l of the detection antigen and 20 μ l of sample were incubated for 9 min. Magnetic Elecsys® beads (30 μ l were added and incubated for 9 min to adsorb the complex of capture antigen - ant-p53 antibody - detection antigen.

Signal measurement was performed on an Elecsys® e601 analyzer according to standard procedures.

The buffer used for the capture antigen comprises the following components:

| <i>Buffer component</i> | <i>conc</i> |
|---|--------------------|
| Kaliumdihydrogenphosphat | 65 mM |
| Di-Kaliumhydrogenphosphat 1x H ₂ O | 35 mM |
| NaCl | 150 mM |
| N-Methylisothiazolon-HCl | 0.01% |
| Oxy-Pyrion | 0.1% |
| Polydocanol | 0.25% |

pH 7.2

The buffer used for the detection antigen comprises the following components:

| <i>Buffer component</i> | <i>conc</i> |
|--------------------------|-------------|
| MES | 50 mM |
| NaCl | 150 mM |
| N-Methylisothiazolon-HCl | 0.01% |
| Oxy-Pyrron | 0.1% |
| Polydocanol | 0.25% |

pH 5.4

Example 3:

Detection of anti-p53 auto-antibodies in clinical samples

- 5 Samples from 463 patients were analyzed side by side for anti-p53 antibodies using the anti-p53 assay as described in Example 2 and the commercially available assay "MESACUP Anti-p53 TEST" distributed by MEDICAL & BIOLOGICAL LABORATORIES CO., LTD., KDX Nagoya, Japan. The later was used according to the instructions given by the manufacturer.

| | Number of samples | Elecsys | Mesacup |
|---------------------------------|--------------------------|--------------------|--------------------|
| | | Sensitivity | Sensitivity |
| Colorectal cancer (CRC) | 88 | 32,95% | 39,77% |
| | | | |
| | | Spezifität | Spezifität |
| Controls from CRC screening | 43 | 97,60% | 86,05% |
| IBD (infectious bowel diseases) | 128 | 96,90% | 79,69% |
| osteoarthritis | 35 | 100% | 80,00% |
| Rheumatoid arthritis | 27 | 96,30% | 81,48% |
| Liver cirrhosis | 17 | 100% | 70,59% |
| Renal failure | 18 | 100% | 83,33% |
| Autoimmun diseases | 40 | 100% | 90,00% |

- 10 As can be seen from the above table, the newly anti-p53 antibody assay detects a comparable amount or slightly less of anti-p53 positive samples in colorectal cancer patients as compared to the competitor assay.

Strikingly the newly developed assay, however, is by far less prone to "positive" signals in disease entities not known to be associated with antibodies to p53.

As shown in Figure 2, the area under the curve (AUC) is better for the assays developed here as compared to a commercially available kit.

5 Most likely the positive measurements found in patients with, controls from CRC screening, IBD (infectious bowel diseases), osteoarthritis, rheumatoid arthritis, liver cirrhosis, renal failure or autoimmun diseases represent false positive measurements. Only very few false positive samples are seen with the new assay, whereas the state of the art assay very often would lead to a false positive results.

10 Obviously, some of the additional positives found with the competitor assay in the CRC cases shown above quite likely are false positives for anti-p53 competitor assay.

The newly developed assay for the in vitro detection of anti-p53 antibodies as disclosed herein has a very good specificity. Such good specificity e.g. translates
15 into less clinical work-up with patients erroneously found as anti-p53 antibody positive.

Patent Claims

1. An in vitro method for detecting an antibody to p53 (anti-p53 antibody) in a sample, the method comprising:
 - a) incubating a sample to be analyzed with a p53 capture antigen and a p53 detection antigen, whereby a complex comprising the p53 capture antigen the anti-p53 antibody and the p53 detection antigen is formed,
 - b) separating the complex formed in (a) from unbound detection antigen,
 - c) measuring the complex obtained in step (b) via the detection antigen comprised therein, thereby detecting the anti-p53 antibody comprised in the sample.
2. The method according to claim 1,
wherein the p53 capture antigen is capable of binding to the solid phase.
3. The method according to 1 or 2,
wherein the p53 capture antigen is covalently bound to a first partner of a binding pair and wherein the second partner of said binding pair is bound to the solid phase.
4. The method according to any of claims 1 to 3,
wherein the first partner of the binding pair is selected from hapten, biotin or biotin analogues such as aminobiotin, iminobiotin or desthiobiotin, DsF and the ligand for a receptor and wherein the second partner of said binding pair is selected from anti-hapten antibody, avidin or streptavidin, FimG DsF, and a ligand-receptor.
5. The method according to any of claims 1 to 4,
wherein the p53 capture antigen is biotinylated and wherein the solid phase is coated with avidin or streptavidin.
6. The method according to claim 1 or 2,
wherein the p53 capture antigen is bound to the solid phase.

7. The method according to any of claims 1 to 6,
wherein the p53 detection antigen comprises a directly detectable label.
8. The method according to any of claims 1 to 7,
wherein the directly detectable label is selected from the group consisting of
5 a luminescent label, a fluorescent label, a chemiluminescent label or an
electrochemiluminescent label.
9. The method according to any of claims 1 to 8,
wherein the directly detectable label is a chemiluminescent or an
electrochemiluminescent label.
10. The method according to any of claims 1 to 9,
wherein both the p53 capture antigen and the p53 detection antigen
comprise a peptide of SEQ ID NO: 1.
11. The method according to any of claims 1 to 10,
wherein both the p53 capture antigen and the p53 detection antigen
15 comprise a peptide of SEQ ID NO: 2.
12. The method according to any of claims 1 to 11,
wherein the peptides of SEQ ID NO:1 and of SEQ ID NO:2 are each
comprised in both the p53 capture antigen and the p53 detection antigen.
13. The method according to any of claims 1 to 12,
wherein the p53 detection antigen comprises both the peptides of SEQ ID
20 NO: 1 and of SEQ ID NO:2 at least twice .
14. The method according to any of claims 1 to 13,
wherein the detection antigen comprises a fusion polypeptide comprising at
least one multimerization domain, at least one polypeptide of SEQ ID NO:
25 1 and at least one polypeptide of SEQ ID NO:2, wherein the
multimerization domain is a prokaryotic or eukaryotic chaperone selected

from the group consisting of FkpA, Skp, SecB, Hsp25, MIP, GroEL, ClpB and ClpX.

15. The method according to any of claims 1 to 14,

wherein the p53 detection antigen comprises an indirectly detectable label.

5 16. A fusion polypeptide comprising at least one multimerization domain at least one polypeptide of SEQ ID NO: 1 and at least one polypeptide of SEQ ID NO:2, wherein the multimerization domain is a prokaryotic or eukaryotic chaperone selected from the group consisting of FkpA, Skp, SecB, Hsp25, MIP, GroEL, ClpB and ClpX.

Figure 1

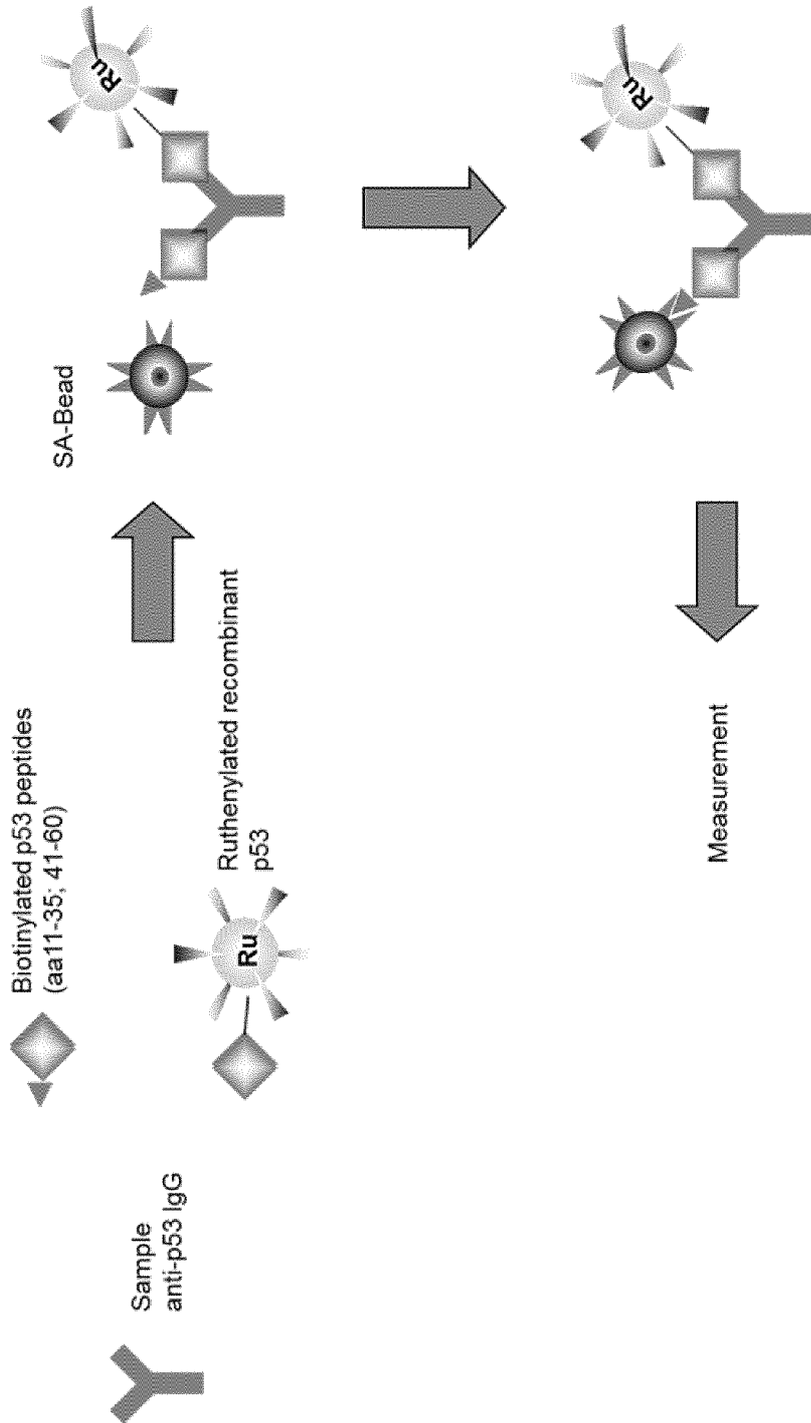
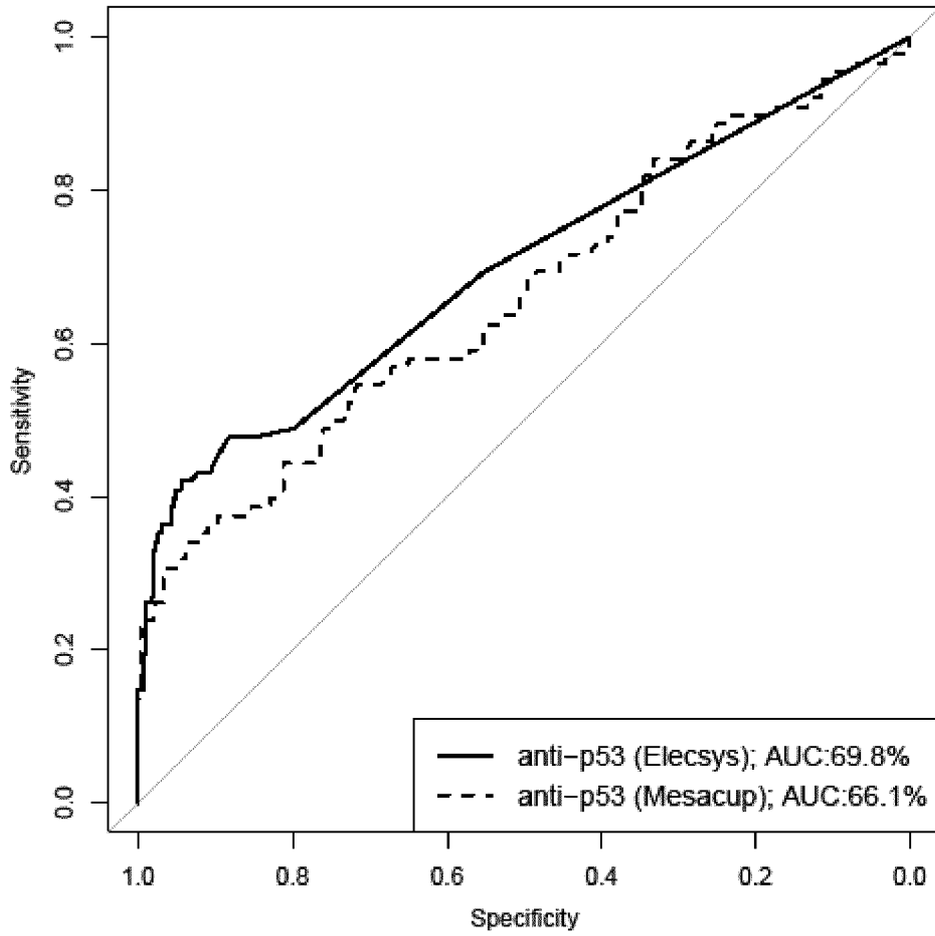
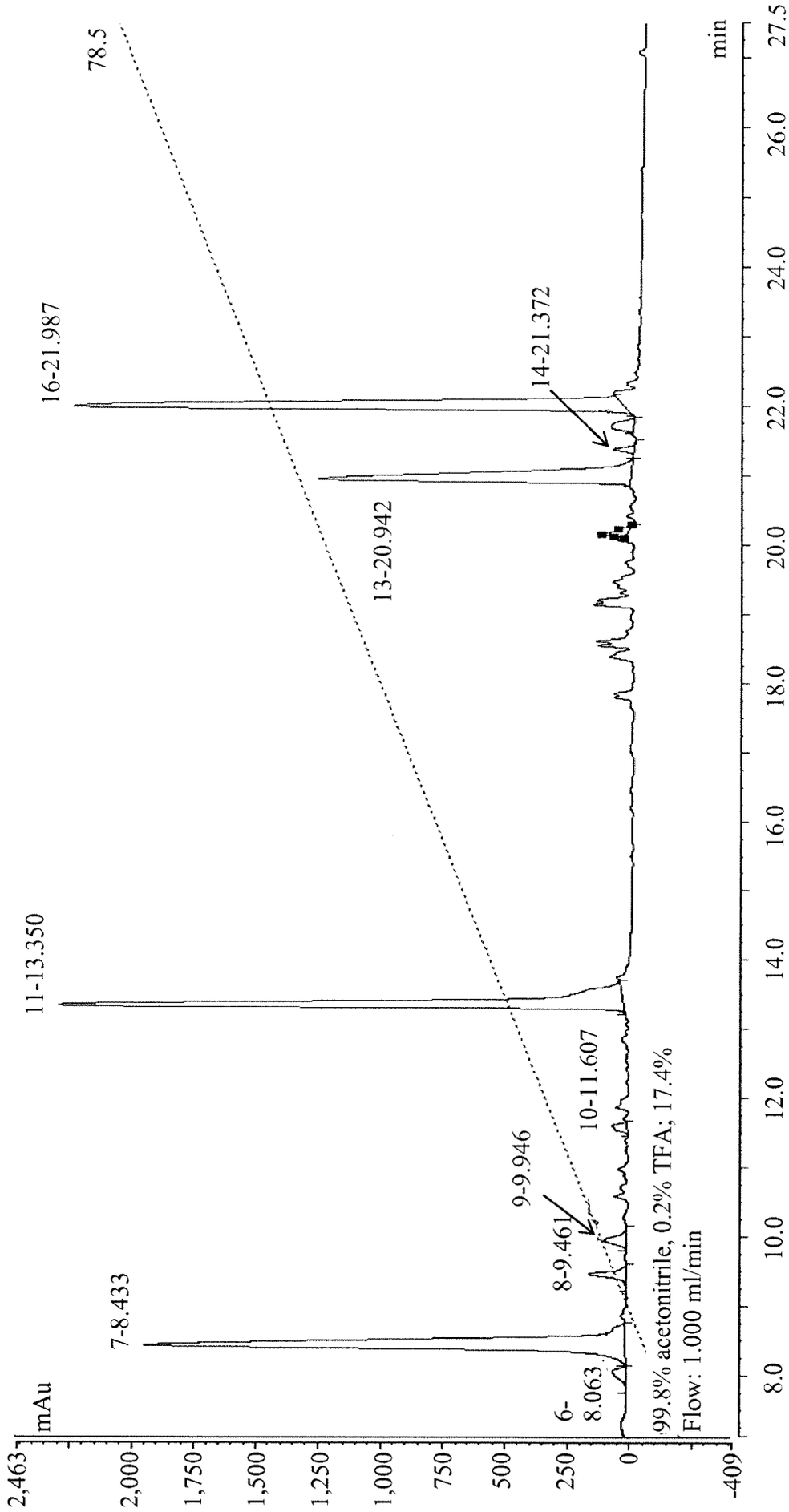


Figure 2





INTERNATIONAL SEARCH REPORT

International application No
PCT/EP2017/055190

A. CLASSIFICATION OF SUBJECT MATTER
INV. G01N33/574 C07K14/47
ADD.
According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED
Minimum documentation searched (classification system followed by classification symbols)
G01N C07K

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)
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Further documents are listed in the continuation of Box C.

See patent family annex.

* Special categories of cited documents :

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| Date of the actual completion of the international search 13 April 2017 | Date of mailing of the international search report 25/04/2017 |
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INTERNATIONAL SEARCH REPORT

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