CPC COOPERATIVE PATENT CLASSIFICATION

C CHEMISTRY; METALLURGY

(NOTES omitted)

CHEMISTRY

C12 BIOCHEMISTRY; BEER; SPIRITS; WINE; VINEGAR; MICROBIOLOGY; ENZYMOLOGY; MUTATION OR GENETIC ENGINEERING (NOTES omitted)

C12Q MEASURING OR TESTING PROCESSES INVOLVING ENZYMES, NUCLEIC ACIDS OR MICROORGANISMS (immunoassay G01N 33/53); COMPOSITIONS OR TEST PAPERS THEREFOR; PROCESSES OF PREPARING SUCH COMPOSITIONS; CONDITION-RESPONSIVE CONTROL IN MICROBIOLOGICAL OR ENZYMOLOGICAL PROCESSES

NOTES

- 1. This subclass <u>does not cover</u> the observation of the progress or of the result of processes specified in this subclass by any of the methods specified in groups <u>G01N 3/00</u> <u>G01N 29/00</u>, which is covered by subclass <u>G01N</u>.
- 2. In this subclass, the following expression is used with the meaning indicated: "involving", when used in relation to a substance, includes the testing for the substance as well as employing the substance as a determinant or reactant in a test for a different substance.
- 3. Attention is drawn to Notes (1) to (3) following the title of class C12.

Measuring or testing processes involving enzymes.

- 4. In this subclass, test media are classified in the appropriate group for the relevant test process.
- 5. In this subclass, it is desirable to add the indexing codes of subclass C12R.
- 6. {Documents describing the use of an electrode for analysis of a specific analyte are classified in C12Q 1/001 or subgroups and not according to the last place rule.}
- 7. {Documents relating to new peptides, e.g. enzymes, or new DNA or its corresponding mRNA, encoding for the peptides, and their use in measuring or testing processes are classified in subclass C07K or in group C12N 9/00 according to the peptides, with the appropriate indexing codes relating to their use in diagnostics. However, where the new nucleic acids are principally used in diagnostic processes, e.g. PCR, hybridisation reactions, the documents are also classified in group C12Q 1/68.}
- 8. {In groups C12Q 1/6876 C12Q 1/6895 and C12Q 1/701 C12Q 1/708 it is compulsory to add the indexing codes C12Q 2600/00 C12Q 2600/178 which reflect the use of the product in combination with the virus groups only if the document relates to products.}
- 9. {In this subclass, combination sets [C-Sets] are used. The detailed information about the C-Sets construction and the associated syntax rules is present in the definitions of C12Q.}

WARNING

1/00

In this subclass non-limiting references (in the sense of paragraph 39 of the Guide to the IPC) may still be displayed in the scheme.

1/005

• • {involving specific analytes or enzymes

1/00	wicasuring or testing processes involving enzymes,	1/005	• • (myorving specific analytes of enzymes
	nucleic acids or microorganisms (measuring		(including groups of enzymes, e.g. oxydases;
	or testing apparatus with condition measuring or		C12Q 1/004 takes precedence)}
	sensing means, e.g. colony counters, C12M 1/34);	1/006	• • · {for glucose}
	Compositions therefor; Processes of preparing such compositions	1/007	• {involving isoenzyme profiles (for detection of an individual isoenzyme C12Q 1/25 - C12Q 1/66)}
	NOTE	1/008	• {for determining co-enzymes or co-factors, e.g. NAD, ATP}
	{In this group, C-Sets are used for classification.	1/02	 involving viable microorganisms
	The detailed information about the C-Sets construction and the associated syntax rules are found in the Definitions of C12Q.}	1/025	• • {for testing or evaluating the effect of chemical or biological compounds, e.g. drugs, cosmetics (antimicrobial activity C12Q 1/18)}
1/001 1/002 1/003 1/004	{Enzyme electrodes}{Electrode membranes}{Functionalisation}{mediator-assisted}	1/04	 Determining presence or kind of microorganism; Use of selective media for testing antibiotics or bacteriocides; Compositions containing a chemical indicator therefor {(C12Q 1/6897 takes precedence)}

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1/045	• • {Culture media therefor}	1/6816 characterised by the detection means
1/06	Quantitative determination	(C12Q 1/6804 takes precedence)
1/08	using multifield media	1/6818 involving interaction of two or more labels,
1/10	Enterobacteria	e.g. resonant energy transfer
1/12	Nitrate to nitrite reducing bacteria	1/682 Signal amplification 1/6823 Release of bound markers
1/14	Streptococcus; Staphylococcus	1/6825 Nucleic acid detection involving sensors
1/16	using radioactive material	1/6827 for detection of mutation or polymorphism
1/18	. Testing for antimicrobial activity of a material	1/683 involving restriction enzymes, e.g. restriction
1/20	using multifield media	fragment length polymorphism [RFLP]
1/22	. Testing for sterility conditions	1/6832 Enhancement of hybridisation reaction
1/24	 Methods of sampling, or inoculating or spreading a sample; Methods of physically isolating an 	1/6834 Enzymatic or biochemical coupling of nucleic
	intact microorganisms	acids to a solid phase
1/25	 involving enzymes not classifiable in groups 	1/6837 using probe arrays or probe chips
	<u>C12Q 1/26</u> {- <u>C12Q 1/66</u> }	(C12Q 1/6874 takes precedence)
1/26	involving oxidoreductase	1/6839 Triple helix formation or other higher order
1/28	involving peroxidase	conformations in hybridisation assays
1/30	involving catalase	1/6841 <u>In situ</u> hybridisation
1/32	involving dehydrogenase	1/6844 Nucleic acid amplification reactions
1/34	 involving hydrolase 	1/6846 {Common amplification features}
1/37	involving peptidase or proteinase	1/6848 characterised by the means for preventing
1/40	involving amylase	contamination or increasing the specificity or
1/42	 involving phosphatase 	sensitivity of an amplification reaction
1/44	• involving esterase	1/6851 Quantitative amplification
1/46	involving cholinesterase	1/6853 using modified primers or templates
1/48	 involving transferase 	1/6858 Ligating adaptors
1/485	• • {involving kinase}	1/6858 Allele-specific amplification
1/50	 involving creatine phosphokinase 	1/686 Polymerase chain reaction [PCR] 1/6862 Ligase chain reaction [LCR]
1/52	• • involving transaminase	1/6865 Promoter-based amplification, e.g. nucleic
1/527	 involving lyase 	acid sequence amplification [NASBA], self-
1/533	 involving isomerase 	sustained sequence replication [3SR] or
1/54	 involving glucose or galactose 	transcription-based amplification system [TAS]
1/56	 involving blood clotting factors, e.g. involving 	1/6867 Replicase-based amplification, e.g. using Q-
	thrombin, thromboplastin, fibrinogen	beta replicase
1/58	involving urea or urease	1/6869 Methods for sequencing
1/60	• involving cholesterol	1/6872 involving mass spectrometry
1/61	involving triglycerides	1/6874 involving nucleic acid arrays, e.g. sequencing
1/62	• involving uric acid	by hybridisation
1/64	• Geomicrobiological testing, e.g. for petroleum	1/6876 Nucleic acid products used in the analysis of
1/66	• involving luciferase	nucleic acids, e.g. primers or probes
1/68	involving nucleic acids	1/6879 for sex determination
	<u>NOTES</u>	1/6881 for tissue or cell typing, e.g. human leukocyte
	1. In this group, classification is made according to	antigen [HLA] probes 1/6883 for diseases caused by alterations of genetic
	the most relevant feature irrespective of the last	1/6883 for diseases caused by alterations of genetic material
	place priority rule.	1/6886 for cancer (immunoassay for cancer
	2. {In groups C12Q 1/68 - C12Q 1/6874, and	G01N 33/574)
	C12Q 1/6897, C-Sets are used for classification.	1/6888 for detection or identification of organisms
	The detailed information about the C-Sets	1/689 for bacteria
	construction and the associated syntax rules are	1/6893 for protozoa
	found in the Definitions of C12Q.	1/6895 for plants, fungi or algae
1/6804	Nucleic acid analysis using immunogens	1/6897 involving reporter genes operably linked to
1,000.	(immunoassay G01N 33/53)	promoters
1/6806	Preparing nucleic acids for analysis, e.g.	1/70 . involving virus or bacteriophage {(immunoassay for
	for polymerase chain reaction [PCR] assay	viruses <u>G01N 33/56983</u>)}
	(C12Q 1/6804 takes precedence)	NOTES
1/6809	Methods for determination or identification of	
	nucleic acids involving differential detection	1. {In this group, classification is made according to the most relevant feature irrespective of the
1/6811	Selection methods for production or design	to the most relevant feature irrespective of the last place priority rule.}
	of target specific oligonucleotides or binding	2. {In this group, C-Sets are used for classification.
1/6012	molecules	The detailed information about the C-Sets
1/6813	Hybridisation assays	

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C12Q 1/70			
(continued)	construction and the associated syntax rules are	2334/50	. Indoles
	found in the Definitions of <u>C12Q</u> .}	2334/52	5-Bromo-4-chloro-3-indolyl, i.e. BCI
1/701	• • {Specific hybridization probes}	2334/70	• the product, e.g. phenol, naphthol being diazotised
1/701	• • {Specific hybridization probes} • • • {for retroviruses}		in situ, e.g. with Fast Red
1/702	{Viruses associated with AIDS}	2337/00	N-linked chromogens for determinations of
		2337700	peptidases and proteinases
1/705	 . • {for herpetoviridae, e.g. herpes simplex, varicella zoster} 	2337/10	Anilides
1/706	,	2337/10	Para-Nitroanilides p-NA
	{for hepatitis}	2337/12	Coumarin derivatives
1/707	{non-A, non-B Hepatitis, excluding hepatitis D}	2337/20	• 7-Amino-4-methylcoumarin, i.e. AMC, MCA
1/708	• • · {for papilloma}	2337/24	• 7-Amino-4-trifluoromethylcoumarin, i.e. AFC
1/708	• • • {for papinoma}	2337/24	Naphthyl amides, e.g. beta-NA, 2-NA, 4-methoxy-
3/00	Condition responsive control processes (apparatus	2331/30	beta-naphthylamine, i.e. 4MNA
	therefor C12M 1/36; controlling or regulating in	2337/40	Rhodamine derivatives
	general G05)	2337/50	Indoles
		2337/50	5-Bromo-4-chloro-3-indolyl, i.e. BCI
2304/00	Chemical means of detecting microorganisms	2331132	5-bromo-4-cmoro-3-mdoryi, i.e. ber
2304/10	. DNA staining	2500/00	Analytical methods involving nucleic acids
2304/12	Ethidium		NOTE
2304/13	Propidium		
2304/16	Acridine orange		Indexing codes <u>C12Q 2500/00</u> - <u>C12Q 2565/634</u>
2304/18	Thionin-type dyes, e.g. Azure, Toluidine Blue		are only used as subsequent symbols in C-Sets and
2304/20	. Redox indicators		are not allocated as single symbols. The detailed
2304/22	Resazurin; Resorufin		information about the C-Sets construction and the
2304/24	Tetrazolium; Formazan		associated syntax rules is present in the Definitions of C12Q.
2304/26	Quinone; Quinol		01 <u>C12Q</u> .
2304/40	. Detection of gases	2520/00	Reactions involving nucleic acids
2304/44	Oxygen	2521/00	Describes the secretarian liberal community and the
2304/46	Carbon dioxide	2521/00	Reaction characterised by the enzymatic activity
2304/48	Ammonia or volatile amines	2521/10	Nucleotidyl transfering
2304/60	. Chemiluminescent detection using ATP-luciferin-	2521/101	DNA polymerase
	luciferase system	2521/107	RNA dependent DNA polymerase, (i.e. reverse transcriptase)
2304/80	Electrochemical detection via electrodes in contact	2521/112	Telomerase
	with culture medium	2521/113 2521/119	
2326/00	Chromogens for determinations of oxidoreductase	2521/119	RNA polymerase
2020/00	enzymes	2521/125	Methyl transferase, i.e. methylase Terminal transferase
2326/10	Benzidines		
2326/12	3,3',5,5'-Tetramethylbenzidine, i.e. TMB	2521/30	Phosphoric diester hydrolysing, i.e. nuclease
2326/14	• Ortho-Tolidine, i.e. 3,3'-dimethyl-(1,1'-	2521/301	. Endonuclease
	biphenyl-4,4'-diamine)	2521/307	
		0501/010	Single strand endonuclease
2326/20		2521/313	Type II endonucleases, i.e. cutting outside
2326/20 2326/30	Ortho-Phenylenediamine		Type II endonucleases, i.e. cutting outside recognition site
2326/20 2326/30	Ortho-Phenylenediamine 2,2'-Azinobis (3-ethylbenzothiazoline-6-sulfonic	2521/319	 Type II endonucleases, i.e. cutting outside recognition site Exonuclease
	Ortho-Phenylenediamine 2,2'-Azinobis (3-ethylbenzothiazoline-6-sulfonic acid), i.e. ABTS	2521/319 2521/325	 Type II endonucleases, i.e. cutting outside recognition site Exonuclease Single stranded exonuclease
2326/30	Ortho-Phenylenediamine 2,2'-Azinobis (3-ethylbenzothiazoline-6-sulfonic	2521/319 2521/325 2521/327	 Type II endonucleases, i.e. cutting outside recognition site Exonuclease Single stranded exonuclease RNAse, e.g. RNAseH
2326/30	Ortho-Phenylenediamine 2,2'-Azinobis (3-ethylbenzothiazoline-6-sulfonic acid), i.e. ABTS 3-Methyl-2-benzothiazolinone hydrazone hydrochloride hydrate, i.e. MBTH	2521/319 2521/325 2521/327 2521/331	 Type II endonucleases, i.e. cutting outside recognition site Exonuclease Single stranded exonuclease RNAse, e.g. RNAseH Methylation site specific nuclease
2326/30 2326/32	 Ortho-Phenylenediamine 2,2'-Azinobis (3-ethylbenzothiazoline-6-sulfonic acid), i.e. ABTS 3-Methyl-2-benzothiazolinone hydrazone 	2521/319 2521/325 2521/327 2521/331 2521/337	 Type II endonucleases, i.e. cutting outside recognition site Exonuclease Single stranded exonuclease RNAse, e.g. RNAseH Methylation site specific nuclease Ribozyme
2326/30 2326/32	 Ortho-Phenylenediamine 2,2'-Azinobis (3-ethylbenzothiazoline-6-sulfonic acid), i.e. ABTS 3-Methyl-2-benzothiazolinone hydrazone hydrochloride hydrate, i.e. MBTH Triphenylmethane dye chromogens, e.g. fluorescein derivatives 	2521/319 2521/325 2521/327 2521/331 2521/337 2521/343	 Type II endonucleases, i.e. cutting outside recognition site Exonuclease Single stranded exonuclease RNAse, e.g. RNAseH Methylation site specific nuclease Ribozyme Abzyme
2326/30 2326/32 2326/40	 Ortho-Phenylenediamine 2,2'-Azinobis (3-ethylbenzothiazoline-6-sulfonic acid), i.e. ABTS 3-Methyl-2-benzothiazolinone hydrazone hydrochloride hydrate, i.e. MBTH Triphenylmethane dye chromogens, e.g. fluorescein 	2521/319 2521/325 2521/327 2521/331 2521/337 2521/343 2521/345	 Type II endonucleases, i.e. cutting outside recognition site Exonuclease Single stranded exonuclease RNAse, e.g. RNAseH Methylation site specific nuclease Ribozyme Abzyme DNAzyme
2326/30 2326/32 2326/40 2326/50	 Ortho-Phenylenediamine 2,2'-Azinobis (3-ethylbenzothiazoline-6-sulfonic acid), i.e. ABTS 3-Methyl-2-benzothiazolinone hydrazone hydrochloride hydrate, i.e. MBTH Triphenylmethane dye chromogens, e.g. fluorescein derivatives Phenols; Naphthols; Catechols Developer 	2521/319 2521/325 2521/327 2521/331 2521/343 2521/345 2521/50	 Type II endonucleases, i.e. cutting outside recognition site Exonuclease Single stranded exonuclease RNAse, e.g. RNAseH Methylation site specific nuclease Ribozyme Abzyme DNAzyme Other enzymatic activities
2326/30 2326/32 2326/40 2326/50 2326/90	 Ortho-Phenylenediamine 2,2'-Azinobis (3-ethylbenzothiazoline-6-sulfonic acid), i.e. ABTS 3-Methyl-2-benzothiazolinone hydrazone hydrochloride hydrate, i.e. MBTH Triphenylmethane dye chromogens, e.g. fluorescein derivatives Phenols; Naphthols; Catechols 	2521/319 2521/325 2521/327 2521/331 2521/337 2521/343 2521/345 2521/50 2521/501	 Type II endonucleases, i.e. cutting outside recognition site Exonuclease Single stranded exonuclease RNAse, e.g. RNAseH Methylation site specific nuclease Ribozyme Abzyme DNAzyme Other enzymatic activities Ligase
2326/30 2326/32 2326/40 2326/50 2326/90 2326/92 2326/96	 Ortho-Phenylenediamine 2,2'-Azinobis (3-ethylbenzothiazoline-6-sulfonic acid), i.e. ABTS 3-Methyl-2-benzothiazolinone hydrazone hydrochloride hydrate, i.e. MBTH Triphenylmethane dye chromogens, e.g. fluorescein derivatives Phenols; Naphthols; Catechols Developer Nitro blue tetrazolium chloride, i.e. NBT 4-Amino-antipyrine 	2521/319 2521/325 2521/327 2521/331 2521/337 2521/343 2521/345 2521/50 2521/501 2521/507	 Type II endonucleases, i.e. cutting outside recognition site Exonuclease Single stranded exonuclease RNAse, e.g. RNAseH Methylation site specific nuclease Ribozyme Abzyme DNAzyme Other enzymatic activities Ligase Recombinase
2326/30 2326/32 2326/40 2326/50 2326/90 2326/92	 Ortho-Phenylenediamine 2,2'-Azinobis (3-ethylbenzothiazoline-6-sulfonic acid), i.e. ABTS 3-Methyl-2-benzothiazolinone hydrazone hydrochloride hydrate, i.e. MBTH Triphenylmethane dye chromogens, e.g. fluorescein derivatives Phenols; Naphthols; Catechols Developer Nitro blue tetrazolium chloride, i.e. NBT 4-Amino-antipyrine O-linked chromogens for determinations of 	2521/319 2521/325 2521/327 2521/331 2521/337 2521/343 2521/345 2521/50 2521/501 2521/507 2521/513	 Type II endonucleases, i.e. cutting outside recognition site Exonuclease Single stranded exonuclease RNAse, e.g. RNAseH Methylation site specific nuclease Ribozyme Abzyme DNAzyme Other enzymatic activities Ligase Recombinase Winding/unwinding enzyme, e.g. helicase
2326/30 2326/32 2326/40 2326/50 2326/90 2326/92 2326/96	 Ortho-Phenylenediamine 2,2'-Azinobis (3-ethylbenzothiazoline-6-sulfonic acid), i.e. ABTS 3-Methyl-2-benzothiazolinone hydrazone hydrochloride hydrate, i.e. MBTH Triphenylmethane dye chromogens, e.g. fluorescein derivatives Phenols; Naphthols; Catechols Developer Nitro blue tetrazolium chloride, i.e. NBT 4-Amino-antipyrine O-linked chromogens for determinations of hydrolase enzymes, e.g. glycosidases, phosphatases, 	2521/319 2521/325 2521/327 2521/331 2521/337 2521/343 2521/345 2521/50 2521/501 2521/507 2521/513 2521/514	 Type II endonucleases, i.e. cutting outside recognition site Exonuclease Single stranded exonuclease RNAse, e.g. RNAseH Methylation site specific nuclease Ribozyme Abzyme DNAzyme Other enzymatic activities Ligase Recombinase Winding/unwinding enzyme, e.g. helicase Mismatch repair protein
2326/30 2326/32 2326/40 2326/50 2326/90 2326/92 2326/96 2334/00	 Ortho-Phenylenediamine 2,2'-Azinobis (3-ethylbenzothiazoline-6-sulfonic acid), i.e. ABTS 3-Methyl-2-benzothiazolinone hydrazone hydrochloride hydrate, i.e. MBTH Triphenylmethane dye chromogens, e.g. fluorescein derivatives Phenols; Naphthols; Catechols Developer Nitro blue tetrazolium chloride, i.e. NBT 4-Amino-antipyrine O-linked chromogens for determinations of hydrolase enzymes, e.g. glycosidases, phosphatases, esterases 	2521/319 2521/325 2521/327 2521/331 2521/337 2521/343 2521/345 2521/50 2521/501 2521/507 2521/513 2521/514 2521/519	 Type II endonucleases, i.e. cutting outside recognition site Exonuclease Single stranded exonuclease RNAse, e.g. RNAseH Methylation site specific nuclease Ribozyme Abzyme DNAzyme Other enzymatic activities Ligase Recombinase Winding/unwinding enzyme, e.g. helicase Mismatch repair protein Topoisomerase
2326/30 2326/32 2326/40 2326/50 2326/90 2326/92 2326/96 2334/00	 Ortho-Phenylenediamine 2,2'-Azinobis (3-ethylbenzothiazoline-6-sulfonic acid), i.e. ABTS 3-Methyl-2-benzothiazolinone hydrazone hydrochloride hydrate, i.e. MBTH Triphenylmethane dye chromogens, e.g. fluorescein derivatives Phenols; Naphthols; Catechols Developer Nitro blue tetrazolium chloride, i.e. NBT 4-Amino-antipyrine O-linked chromogens for determinations of hydrolase enzymes, e.g. glycosidases, phosphatases, esterases p-Nitrophenol derivatives 	2521/319 2521/325 2521/327 2521/331 2521/343 2521/345 2521/50 2521/501 2521/507 2521/513 2521/514 2521/519 2521/525	 Type II endonucleases, i.e. cutting outside recognition site Exonuclease Single stranded exonuclease RNAse, e.g. RNAseH Methylation site specific nuclease Ribozyme Abzyme DNAzyme Other enzymatic activities Ligase Recombinase Winding/unwinding enzyme, e.g. helicase Mismatch repair protein Topoisomerase Phosphatase
2326/30 2326/32 2326/40 2326/50 2326/90 2326/92 2326/96 2334/00 2334/10 2334/20	 Ortho-Phenylenediamine 2,2'-Azinobis (3-ethylbenzothiazoline-6-sulfonic acid), i.e. ABTS 3-Methyl-2-benzothiazolinone hydrazone hydrochloride hydrate, i.e. MBTH Triphenylmethane dye chromogens, e.g. fluorescein derivatives Phenols; Naphthols; Catechols Developer Nitro blue tetrazolium chloride, i.e. NBT 4-Amino-antipyrine O-linked chromogens for determinations of hydrolase enzymes, e.g. glycosidases, phosphatases, esterases p-Nitrophenol derivatives Coumarin derivatives 	2521/319 2521/325 2521/327 2521/331 2521/337 2521/343 2521/50 2521/501 2521/507 2521/513 2521/514 2521/519 2521/525 2521/531	 Type II endonucleases, i.e. cutting outside recognition site Exonuclease Single stranded exonuclease RNAse, e.g. RNAseH Methylation site specific nuclease Ribozyme Abzyme DNAzyme Other enzymatic activities Ligase Recombinase Winding/unwinding enzyme, e.g. helicase Mismatch repair protein Topoisomerase Phosphatase Glycosylase
2326/30 2326/32 2326/40 2326/50 2326/90 2326/92 2326/96 2334/00	 Ortho-Phenylenediamine 2,2'-Azinobis (3-ethylbenzothiazoline-6-sulfonic acid), i.e. ABTS 3-Methyl-2-benzothiazolinone hydrazone hydrochloride hydrate, i.e. MBTH Triphenylmethane dye chromogens, e.g. fluorescein derivatives Phenols; Naphthols; Catechols Developer Nitro blue tetrazolium chloride, i.e. NBT 4-Amino-antipyrine O-linked chromogens for determinations of hydrolase enzymes, e.g. glycosidases, phosphatases, esterases p-Nitrophenol derivatives Coumarin derivatives 4-Methylumbelliferyl, i.e. beta- 	2521/319 2521/325 2521/327 2521/331 2521/337 2521/343 2521/345 2521/501 2521/507 2521/507 2521/513 2521/514 2521/519 2521/525 2521/531 2521/537	 Type II endonucleases, i.e. cutting outside recognition site Exonuclease Single stranded exonuclease RNAse, e.g. RNAseH Methylation site specific nuclease Ribozyme Abzyme DNAzyme Other enzymatic activities Ligase Recombinase Winding/unwinding enzyme, e.g. helicase Mismatch repair protein Topoisomerase Phosphatase Glycosylase Protease
2326/30 2326/32 2326/40 2326/50 2326/90 2326/92 2326/96 2334/00 2334/10 2334/20 2334/20	 Ortho-Phenylenediamine 2,2'-Azinobis (3-ethylbenzothiazoline-6-sulfonic acid), i.e. ABTS 3-Methyl-2-benzothiazolinone hydrazone hydrochloride hydrate, i.e. MBTH Triphenylmethane dye chromogens, e.g. fluorescein derivatives Phenols; Naphthols; Catechols Developer Nitro blue tetrazolium chloride, i.e. NBT 4-Amino-antipyrine O-linked chromogens for determinations of hydrolase enzymes, e.g. glycosidases, phosphatases, esterases p-Nitrophenol derivatives Coumarin derivatives 4-Methylumbelliferyl, i.e. betamethylumbelliferone, 4MU 	2521/319 2521/325 2521/327 2521/331 2521/337 2521/345 2521/50 2521/501 2521/507 2521/513 2521/514 2521/519 2521/525 2521/531 2521/537 2521/539	 Type II endonucleases, i.e. cutting outside recognition site Exonuclease Single stranded exonuclease RNAse, e.g. RNAseH Methylation site specific nuclease Ribozyme Abzyme DNAzyme Other enzymatic activities Ligase Recombinase Winding/unwinding enzyme, e.g. helicase Mismatch repair protein Topoisomerase Phosphatase Glycosylase Protease Deaminase
2326/30 2326/32 2326/40 2326/50 2326/90 2326/92 2326/96 2334/00 2334/10 2334/20	 Ortho-Phenylenediamine 2,2'-Azinobis (3-ethylbenzothiazoline-6-sulfonic acid), i.e. ABTS 3-Methyl-2-benzothiazolinone hydrazone hydrochloride hydrate, i.e. MBTH Triphenylmethane dye chromogens, e.g. fluorescein derivatives Phenols; Naphthols; Catechols Developer Nitro blue tetrazolium chloride, i.e. NBT 4-Amino-antipyrine O-linked chromogens for determinations of hydrolase enzymes, e.g. glycosidases, phosphatases, esterases p-Nitrophenol derivatives Coumarin derivatives 4-Methylumbelliferyl, i.e. betamethylumbelliferone, 4MU Naphthol derivatives, e.g. alpha-naphthyl-esters, i.e. 	2521/319 2521/325 2521/327 2521/331 2521/337 2521/343 2521/345 2521/501 2521/507 2521/507 2521/513 2521/514 2521/519 2521/525 2521/531 2521/537	 Type II endonucleases, i.e. cutting outside recognition site Exonuclease Single stranded exonuclease RNAse, e.g. RNAseH Methylation site specific nuclease Ribozyme Abzyme DNAzyme Other enzymatic activities Ligase Recombinase Winding/unwinding enzyme, e.g. helicase Mismatch repair protein Topoisomerase Phosphatase Glycosylase Protease
2326/30 2326/32 2326/40 2326/50 2326/90 2326/92 2326/96 2334/00 2334/10 2334/20 2334/22 2334/30	 Ortho-Phenylenediamine 2,2'-Azinobis (3-ethylbenzothiazoline-6-sulfonic acid), i.e. ABTS 3-Methyl-2-benzothiazolinone hydrazone hydrochloride hydrate, i.e. MBTH Triphenylmethane dye chromogens, e.g. fluorescein derivatives Phenols; Naphthols; Catechols Developer Nitro blue tetrazolium chloride, i.e. NBT 4-Amino-antipyrine O-linked chromogens for determinations of hydrolase enzymes, e.g. glycosidases, phosphatases, esterases p-Nitrophenol derivatives Coumarin derivatives 4-Methylumbelliferyl, i.e. betamethylumbelliferone, 4MU Naphthol derivatives, e.g. alpha-naphthyl-esters, i.e. alpha-NE, beta-naphthyl-esters, i.e. beta-NE 	2521/319 2521/325 2521/327 2521/331 2521/337 2521/343 2521/50 2521/501 2521/507 2521/513 2521/514 2521/519 2521/525 2521/531 2521/537 2521/539 2521/543	 Type II endonucleases, i.e. cutting outside recognition site Exonuclease Single stranded exonuclease RNAse, e.g. RNAseH Methylation site specific nuclease Ribozyme Abzyme DNAzyme Other enzymatic activities Ligase Recombinase Winding/unwinding enzyme, e.g. helicase Mismatch repair protein Topoisomerase Phosphatase Glycosylase Protease Deaminase Immobilised enzyme(s)
2326/30 2326/32 2326/40 2326/50 2326/90 2326/92 2326/96 2334/00 2334/10 2334/20 2334/20	 Ortho-Phenylenediamine 2,2'-Azinobis (3-ethylbenzothiazoline-6-sulfonic acid), i.e. ABTS 3-Methyl-2-benzothiazolinone hydrazone hydrochloride hydrate, i.e. MBTH Triphenylmethane dye chromogens, e.g. fluorescein derivatives Phenols; Naphthols; Catechols Developer Nitro blue tetrazolium chloride, i.e. NBT 4-Amino-antipyrine O-linked chromogens for determinations of hydrolase enzymes, e.g. glycosidases, phosphatases, esterases p-Nitrophenol derivatives Coumarin derivatives 4-Methylumbelliferyl, i.e. betamethylumbelliferone, 4MU Naphthol derivatives, e.g. alpha-naphthyl-esters, i.e. 	2521/319 2521/325 2521/327 2521/331 2521/337 2521/345 2521/50 2521/501 2521/507 2521/513 2521/514 2521/519 2521/525 2521/531 2521/537 2521/539	 Type II endonucleases, i.e. cutting outside recognition site Exonuclease Single stranded exonuclease RNAse, e.g. RNAseH Methylation site specific nuclease Ribozyme Abzyme DNAzyme Other enzymatic activities Ligase Recombinase Winding/unwinding enzyme, e.g. helicase Mismatch repair protein Topoisomerase Phosphatase Glycosylase Protease Deaminase

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2522/101	Nucleic acid binding proteins	2525/205	Aptamer
2522/101	Single or double stranded nucleic acid binding	2525/207	siRNA, miRNA
	proteins	2525/30	Oligonucleotides characterised by their secondary
2523/00	Reactions characterised by treatment of reaction	2525/201	structure
	samples	2525/301	Hairpin oligonucleotides
2523/10	Characterised by chemical treatment	2525/307	. Circular oligonucleotides
2523/101	Crosslinking agents, e.g. psoralen	2525/313	Branched oligonucleotides
2523/107	Chemical cleaving agents	2527/00	Reactions demanding special reaction conditions
2523/109	chemical ligation between nucleic acids	2527/101	. Temperature
2523/113	Denaturating agents	2527/107	• Temperature of melting, i.e. Tm
2523/115	• • oxidising agents	2527/109	• Pressure
2523/119	Renaturing agents	2527/113	. Time
2523/125	Bisulfite(s)	2527/119	. pH
2523/30	Characterised by physical treatment	2527/125	Specific component of sample, medium or buffer
2523/301	Sonication	2527/127	the enzyme inhibitor or activator used
2523/303	Applying a physical force on a nucleic acid	2527/137	Concentration of a component of medium
2523/305	Denaturation or renaturation by physical action	2527/143	Concentration of primer or probe
2523/307	Denaturation or renaturation by electric current/	2527/146	Concentration of target or template
7500/200	voltage	2527/149	Concentration of an enzyme
2523/308	. Adsorption or desorption	2527/15	. Gradients
2523/31	Electrostatic interactions, e.g. use of cationic polymers in hybridisation reactions	2527/153	. Viscosity
2523/313	Irradiation, e.g. UV irradiation	2527/156	. Permeability
2523/319	Photocleavage, photolysis, photoactivation	2531/00	Reactions of nucleic acids characterised by
2523/31	Centrifugation	2531/10	• the purpose being amplify/increase the copy number
25 25 1 5 2			of target nucleic acid
2525/00	Reactions involving modified oligonucleotides,	2531/101	. Linear amplification, i.e. non exponential
	nucleic acids, or nucleotides	2531/107	• • Probe or oligonucleotide ligation
525/10	Modifications characterised by	2531/113	PCR
2525/101	incorporating non-naturally occurring	2531/119	Strand displacement amplification [SDA]
505/107	nucleotides, e.g. inosine	2531/125	Rolling circle
2525/107 2525/113	incorporating a peptide nucleic acid	2531/131	Inverse PCR
2525/115	incorporating modified backbone	2531/137	Ligase Chain Reaction [LCR]
2525/117	incorporating modified baseincorporating abasic sites	2531/143	Promoter based amplification, e.g. NASBA, 3SR
2525/119	incorporating abasic sites incorporating both deoxyribonucleotides and		TAS
	ribonucleotides	2531/149	Replicase based amplification, e.g. Q beta replicase
	incorporating agents resulting in resistance to	2533/00	Describeration of the second s
2525/125			
	degradation	2555/00	Reactions characterised by the enzymatic reaction
525/131	incorporating a restriction site		principle used
2525/131		2533/10	 principle used the purpose being to increase the length of an oligonucleotide strand
2525/131 2525/137	incorporating a restriction siteincorporating/modifying moieties to eliminate	2533/10 2533/101	 principle used the purpose being to increase the length of an oligonucleotide strand Primer extension
2525/131 2525/137 2525/143	 incorporating a restriction site incorporating/modifying moieties to eliminate restriction sites incorporating a promoter sequence incorporating a coding sequence 	2533/10	 principle used the purpose being to increase the length of an oligonucleotide strand
2.525/131 2.525/137 2.525/143 2.525/149	 incorporating a restriction site incorporating/modifying moieties to eliminate restriction sites incorporating a promoter sequence incorporating a coding sequence incorporating a consensus or conserved sequence 	2533/10 2533/101	 principle used the purpose being to increase the length of an oligonucleotide strand Primer extension Probe or oligonucleotide ligation
525/131 525/137 525/143 525/149 525/15	 incorporating a restriction site incorporating/modifying moieties to eliminate restriction sites incorporating a promoter sequence incorporating a coding sequence incorporating a consensus or conserved sequence repeat or repeated sequences, e.g. VNTR, 	2533/10 2533/101 2533/107	 principle used the purpose being to increase the length of an oligonucleotide strand Primer extension
2.525/131 2.525/137 2.525/143 2.525/149 2.525/15 2.525/151	 incorporating a restriction site incorporating/modifying moieties to eliminate restriction sites incorporating a promoter sequence incorporating a coding sequence incorporating a consensus or conserved sequence repeat or repeated sequences, e.g. VNTR, microsatellite, concatemer 	2533/10 2533/101 2533/107	 principle used the purpose being to increase the length of an oligonucleotide strand Primer extension Probe or oligonucleotide ligation Reactions characterised by the assay type for
2.525/131 2.525/137 2.525/143 2.525/149 2.525/15 2.525/151	 incorporating a restriction site incorporating/modifying moieties to eliminate restriction sites incorporating a promoter sequence incorporating a coding sequence incorporating a consensus or conserved sequence repeat or repeated sequences, e.g. VNTR, microsatellite, concatemer incorporating/generating a new priming site 	2533/10 2533/101 2533/107	 principle used the purpose being to increase the length of an oligonucleotide strand Primer extension Probe or oligonucleotide ligation Reactions characterised by the assay type for determining the identity of a nucleotide base or a sequence of oligonucleotides Sanger sequencing method, i.e. oligonucleotide
2.525/131 2.525/137 2.525/143 2.525/149 2.525/15 2.525/151	 incorporating a restriction site incorporating/modifying moieties to eliminate restriction sites incorporating a promoter sequence incorporating a coding sequence incorporating a consensus or conserved sequence repeat or repeated sequences, e.g. VNTR, microsatellite, concatemer 	2533/101 2533/101 2533/107 2535/00	 principle used the purpose being to increase the length of an oligonucleotide strand Primer extension Probe or oligonucleotide ligation Reactions characterised by the assay type for determining the identity of a nucleotide base or a sequence of oligonucleotides Sanger sequencing method, i.e. oligonucleotide sequencing using primer elongation and
2.525/131 2.525/137 2.525/143 2.525/149 2.525/15 2.525/151 2.525/151 2.525/161	 incorporating a restriction site incorporating/modifying moieties to eliminate restriction sites incorporating a promoter sequence incorporating a coding sequence incorporating a consensus or conserved sequence repeat or repeated sequences, e.g. VNTR, microsatellite, concatemer incorporating/generating a new priming site incorporating target specific and non-target specific sites incorporating a polynucleotide run, e.g. polyAs, 	2533/101 2533/101 2533/107 2535/00	 principle used the purpose being to increase the length of an oligonucleotide strand Primer extension Probe or oligonucleotide ligation Reactions characterised by the assay type for determining the identity of a nucleotide base or a sequence of oligonucleotides Sanger sequencing method, i.e. oligonucleotide sequencing using primer elongation and dideoxynucleotides as chain terminators Maxam and Gilbert method, i.e. sequential release
525/131 525/137 525/143 525/149 525/15 525/151 525/151 525/151 525/161	 incorporating a restriction site incorporating/modifying moieties to eliminate restriction sites incorporating a promoter sequence incorporating a coding sequence incorporating a consensus or conserved sequence repeat or repeated sequences, e.g. VNTR, microsatellite, concatemer incorporating/generating a new priming site incorporating target specific and non-target specific sites incorporating a polynucleotide run, e.g. polyAs, polyTs 	2533/10 2533/101 2533/107 2535/00 2535/101 2535/107	 principle used the purpose being to increase the length of an oligonucleotide strand Primer extension Probe or oligonucleotide ligation Reactions characterised by the assay type for determining the identity of a nucleotide base or a sequence of oligonucleotides Sanger sequencing method, i.e. oligonucleotide sequencing using primer elongation and dideoxynucleotides as chain terminators Maxam and Gilbert method, i.e. sequential release and detection of nucleotides
525/131 525/137 525/143 525/149 525/15 525/151 525/151 525/153 525/161	 incorporating a restriction site incorporating/modifying moieties to eliminate restriction sites incorporating a promoter sequence incorporating a coding sequence incorporating a consensus or conserved sequence repeat or repeated sequences, e.g. VNTR, microsatellite, concatemer incorporating/generating a new priming site incorporating target specific and non-target specific sites incorporating a polynucleotide run, e.g. polyAs, polyTs incorporating arbitrary or random nucleotide 	2533/10 2533/101 2533/107 2535/00 2535/101 2535/107 2535/113	 principle used the purpose being to increase the length of an oligonucleotide strand Primer extension Probe or oligonucleotide ligation Reactions characterised by the assay type for determining the identity of a nucleotide base or a sequence of oligonucleotides Sanger sequencing method, i.e. oligonucleotide sequencing using primer elongation and dideoxynucleotides as chain terminators Maxam and Gilbert method, i.e. sequential release and detection of nucleotides Cycle sequencing
525/131 525/137 525/143 525/149 525/15 525/151 525/155 525/161 525/173 525/179	 incorporating a restriction site incorporating/modifying moieties to eliminate restriction sites incorporating a promoter sequence incorporating a coding sequence incorporating a consensus or conserved sequence repeat or repeated sequences, e.g. VNTR, microsatellite, concatemer incorporating/generating a new priming site incorporating target specific and non-target specific sites incorporating a polynucleotide run, e.g. polyAs, polyTs incorporating arbitrary or random nucleotide sequences 	2533/10 2533/101 2533/107 2535/00 2535/101 2535/107 2535/113 2535/119	 principle used the purpose being to increase the length of an oligonucleotide strand Primer extension Probe or oligonucleotide ligation Reactions characterised by the assay type for determining the identity of a nucleotide base or a sequence of oligonucleotides Sanger sequencing method, i.e. oligonucleotide sequencing using primer elongation and dideoxynucleotides as chain terminators Maxam and Gilbert method, i.e. sequential release and detection of nucleotides Cycle sequencing Double strand sequencing
525/131 525/137 525/143 525/149 525/15 525/151 525/155 525/161 525/173 525/179	 incorporating a restriction site incorporating/modifying moieties to eliminate restriction sites incorporating a promoter sequence incorporating a coding sequence incorporating a consensus or conserved sequence repeat or repeated sequences, e.g. VNTR, microsatellite, concatemer incorporating/generating a new priming site incorporating target specific and non-target specific sites incorporating a polynucleotide run, e.g. polyAs, polyTs incorporating arbitrary or random nucleotide sequences incorporating bases where the precise position of 	2533/10 2533/101 2533/107 2535/00 2535/101 2535/107 2535/113 2535/119 2535/122	 principle used the purpose being to increase the length of an oligonucleotide strand Primer extension Probe or oligonucleotide ligation Reactions characterised by the assay type for determining the identity of a nucleotide base or a sequence of oligonucleotides Sanger sequencing method, i.e. oligonucleotide sequencing using primer elongation and dideoxynucleotides as chain terminators Maxam and Gilbert method, i.e. sequential release and detection of nucleotides Cycle sequencing Double strand sequencing Massive parallel sequencing
525/131 525/137 525/143 525/149 525/15 525/151 525/151 525/161 525/173 525/179 525/185	 incorporating a restriction site incorporating/modifying moieties to eliminate restriction sites incorporating a promoter sequence incorporating a coding sequence incorporating a consensus or conserved sequence repeat or repeated sequences, e.g. VNTR, microsatellite, concatemer incorporating/generating a new priming site incorporating target specific and non-target specific sites incorporating a polynucleotide run, e.g. polyAs, polyTs incorporating arbitrary or random nucleotide sequences incorporating bases where the precise position of the bases in the nucleic acid string is important 	2533/10 2533/101 2533/107 2535/00 2535/101 2535/107 2535/113 2535/119 2535/122 2535/125	 principle used the purpose being to increase the length of an oligonucleotide strand Primer extension Probe or oligonucleotide ligation Reactions characterised by the assay type for determining the identity of a nucleotide base or a sequence of oligonucleotides Sanger sequencing method, i.e. oligonucleotide sequencing using primer elongation and dideoxynucleotides as chain terminators Maxam and Gilbert method, i.e. sequential release and detection of nucleotides Cycle sequencing Double strand sequencing Massive parallel sequencing Allele specific primer extension
525/131 525/137 525/143 525/149 525/15 525/151 525/151 525/153 525/161 525/173 525/179 525/185	 incorporating a restriction site incorporating/modifying moieties to eliminate restriction sites incorporating a promoter sequence incorporating a coding sequence incorporating a consensus or conserved sequence repeat or repeated sequences, e.g. VNTR, microsatellite, concatemer incorporating/generating a new priming site incorporating target specific and non-target specific sites incorporating a polynucleotide run, e.g. polyAs, polyTs incorporating arbitrary or random nucleotide sequences incorporating bases where the precise position of the bases in the nucleic acid string is important incorporating a non-extendable or blocking 	2533/10 2533/107 2533/107 2535/00 2535/101 2535/107 2535/113 2535/119 2535/122 2535/125 2535/131	 principle used the purpose being to increase the length of an oligonucleotide strand Primer extension Probe or oligonucleotide ligation Reactions characterised by the assay type for determining the identity of a nucleotide base or a sequence of oligonucleotides Sanger sequencing method, i.e. oligonucleotide sequencing using primer elongation and dideoxynucleotides as chain terminators Maxam and Gilbert method, i.e. sequential release and detection of nucleotides Cycle sequencing Double strand sequencing Massive parallel sequencing Allele specific primer extension Allele specific probes
525/131 525/137 525/143 525/149 525/15 525/151 525/151 525/161 525/173 525/179 525/185 525/186	 incorporating a restriction site incorporating/modifying moieties to eliminate restriction sites incorporating a promoter sequence incorporating a coding sequence incorporating a consensus or conserved sequence repeat or repeated sequences, e.g. VNTR, microsatellite, concatemer incorporating/generating a new priming site incorporating target specific and non-target specific sites incorporating a polynucleotide run, e.g. polyAs, polyTs incorporating arbitrary or random nucleotide sequences incorporating bases where the precise position of the bases in the nucleic acid string is important incorporating a non-extendable or blocking moiety 	2533/10 2533/101 2533/107 2535/00 2535/101 2535/107 2535/113 2535/119 2535/122 2535/125 2535/131 2535/137	 principle used the purpose being to increase the length of an oligonucleotide strand Primer extension Probe or oligonucleotide ligation Reactions characterised by the assay type for determining the identity of a nucleotide base or a sequence of oligonucleotides Sanger sequencing method, i.e. oligonucleotide sequencing using primer elongation and dideoxynucleotides as chain terminators Maxam and Gilbert method, i.e. sequential release and detection of nucleotides Cycle sequencing Double strand sequencing Massive parallel sequencing Allele specific primer extension Allele specific probes Amplification Refractory Mutation System [ARMS]
2525/131 2525/137 2525/143 2525/149 2525/15 2525/151 2525/151 2525/151 2525/161 2525/173 2525/179 2525/185 2525/186	 incorporating a restriction site incorporating/modifying moieties to eliminate restriction sites incorporating a promoter sequence incorporating a coding sequence incorporating a consensus or conserved sequence repeat or repeated sequences, e.g. VNTR, microsatellite, concatemer incorporating/generating a new priming site incorporating target specific and non-target specific sites incorporating a polynucleotide run, e.g. polyAs, polyTs incorporating arbitrary or random nucleotide sequences incorporating bases where the precise position of the bases in the nucleic acid string is important incorporating a non-extendable or blocking 	2533/10 2533/101 2533/107 2535/00 2535/101 2535/107 2535/113 2535/122 2535/125 2535/131 2535/137 2535/138	 principle used the purpose being to increase the length of an oligonucleotide strand Primer extension Probe or oligonucleotide ligation Reactions characterised by the assay type for determining the identity of a nucleotide base or a sequence of oligonucleotides Sanger sequencing method, i.e. oligonucleotide sequencing using primer elongation and dideoxynucleotides as chain terminators Maxam and Gilbert method, i.e. sequential release and detection of nucleotides Cycle sequencing Double strand sequencing Massive parallel sequencing Allele specific primer extension Allele specific probes Amplification Refractory Mutation System [ARMS] Amplified fragment length polymorphism [AFLP]
2525/125 2525/131 2525/137 2525/137 2525/143 2525/149 2525/15 2525/15 2525/151 2525/151 2525/173 2525/179 2525/186 2525/186 2525/191 2525/197 2525/203	 incorporating a restriction site incorporating/modifying moieties to eliminate restriction sites incorporating a promoter sequence incorporating a coding sequence incorporating a consensus or conserved sequence repeat or repeated sequences, e.g. VNTR, microsatellite, concatemer incorporating/generating a new priming site incorporating target specific and non-target specific sites incorporating a polynucleotide run, e.g. polyAs, polyTs incorporating arbitrary or random nucleotide sequences incorporating bases where the precise position of the bases in the nucleic acid string is important incorporating a non-extendable or blocking moiety incorporating an adaptor 	2533/10 2533/101 2533/107 2535/00 2535/101 2535/107 2535/113 2535/119 2535/122 2535/125 2535/131 2535/137	 principle used the purpose being to increase the length of an oligonucleotide strand Primer extension Probe or oligonucleotide ligation Reactions characterised by the assay type for determining the identity of a nucleotide base or a sequence of oligonucleotides Sanger sequencing method, i.e. oligonucleotide sequencing using primer elongation and dideoxynucleotides as chain terminators Maxam and Gilbert method, i.e. sequential release and detection of nucleotides Cycle sequencing Double strand sequencing Massive parallel sequencing Allele specific primer extension Allele specific probes Amplification Refractory Mutation System [ARMS]

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2537/00	Reactions characterised by the reaction format or	2547/107	Use of permeable barriers, e.g. waxes
2527/10	use of a specific feature	2549/00	Reactions characterised by the features used to
2537/10 2537/101	the purpose or use ofHomogeneous assay format, e.g. one pot reaction		influence the efficiency or specificity
2537/107	Homoduplex formation	2549/10	• the purpose being that of reducing false positive or
2537/107	Heteroduplex formation		false negative signals
2537/119	. Triple helix formation	2549/101	Hot start
2537/125	Sandwich assay format	2549/107	Cold start
2537/137	. a displacement step	2549/113	using nested probes
2537/1373	Displacement by a nucleic acid	2549/119	using nested primers
2537/1376	Displacement by an enzyme	2549/125	using sterilising/blocking agents, e.g. albumin
2537/143	Multiplexing, i.e. use of multiple primers	2549/126	using oligonucleotides as clamps
	or probes in a single reaction, usually for simultaneously analyse of multiple analysis	2560/00	Nucleic acid detection
2537/149	Sequential reactions	2561/00	Nucleic acid detection characterised by assay
2537/155	Cyclic reactions	2561/101	method
2537/157	A reaction step characterised by the number of	2561/101	. Taqman
	molecules incorporated or released	2561/107	Enzyme complementation Helpidication approaching according to the property of the proper
2537/159	Reduction of complexity, e.g. amplification of	2561/108	Hybridisation protection assay [HPA]
	subsets, removing duplicated genomic regions	2561/109	Invader technology Productions are a second and a second are a second and a second are a s
2537/16	Assays for determining copy number or wherein	2561/113	Real time assay
	the copy number is of special importance	2561/119	Fluorescence polarisation Fluorescence lifetime massyrement
2537/161	A competitive reaction step	2561/12	Fluorescence lifetime measurement Linear Detection Reportion II DRI
2537/162	Helper probe	2561/125	Ligase Detection Reaction [LDR] Protein transaction access.
2537/163	blocking probe	2561/127	Protein truncation assay
2537/164	Methylation detection other then bisulfite or	2563/00	Nucleic acid detection characterized by the use of
0505/165	methylation sensitive restriction endonucleases		physical, structural and functional properties
2537/165	Mathematical modelling, e.g. logarithm, ratio	2563/101	radioactivity, e.g. radioactive labels
2539/00	Reactions characterised by analysis of gene	2563/103	. luminescence
	expression or genome comparison	2563/107	. fluorescence
2539/10	The purpose being sequence identification by	2563/113	• the label being electroactive, e.g. redox labels
	analysis of gene expression or genome comparison	2563/116	• electrical properties of nucleic acids, e.g.
	characterised by		impedance, conductivity or resistance
2539/101	Subtraction analysis	2563/119	the label being proteinic
2539/103	Serial analysis of gene expression [SAGE]	2563/125	• the label being enzymatic, i.e. proteins, and non
2539/105	Involving introns, exons, or splice junctions		proteins, such as nucleic acid with enzymatic activity
2539/107	Representational Difference Analysis [RDA]	2563/131	• the label being a member of a cognate binding pair,
2539/113	. Differential Display Analysis [DDA]	2303/131	i.e. extends to antibodies, haptens, avidin
2539/115	Comparative genomic hybridisation [CGH]	2563/137	• Metal/ion, e.g. metal label
2541/00	Reactions characterised by directed evolution	2563/143	Magnetism, e.g. magnetic label
2541/10	the purpose being the selection or design of target	2563/149	• Particles, e.g. beads
	specific nucleic acid binding sequences	2563/155	Particles of a defined size, e.g. nanoparticles
2541/101	Selex	2563/157	Nanotubes or nanorods
2543/00	Reactions characterised by the reaction site, e.g.	2563/159	• Microreactors, e.g. emulsion PCR or sequencing,
2343/00	cell or chromosome		droplet PCR, microcapsules, i.e. non-liquid
2543/10	• the purpose being "in situ" analysis		containers with a range of different permeability's
2543/101	in situ amplification		for different reaction components
	-	2563/161	• Vesicles, e.g. liposome
2545/00	Reactions characterised by their quantitative	2563/167	. Mass label
	nature	2563/173	• staining/intercalating agent, e.g. ethidium bromide
2545/10	the purpose being quantitative analysis	2563/179	the label being a nucleic acid
2545/101	with an internal standard/control	2563/185	Nucleic acid dedicated to use as a hidden marker/
2545/107	• with a competitive internal standard/control		bar code, e.g. inclusion of nucleic acids to mark art
2545/113	with an external standard/control, i.e. control		objects or animals
2545/114	reaction is separated from the test/target reaction . involving a quantitation step	2565/00	Nucleic acid analysis characterised by mode or
		05.65.45.0	means of detection
2547/00	Reactions characterised by the features used to	2565/10	Detection mode being characterised by the assay principle
25/7/10	prevent contamination	2565/101	principle Intersection between at least two lebels
2547/10 2547/101	the purpose being preventing contamination by confinement to a single tube/container	2565/101 2565/1015	Interaction between at least two labels labels being on the same oligonucleotide
<i>2341/101</i>	• • by commement to a single tube/container	2565/1015	
		2303/102	• • Multiple non-interacting labels

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2565/1025	labels being on the same oligonucleotide
2565/107	Alteration in the property of hybridised versus
	free label oligonucleotides
2565/113	based on agglutination/precipitation
2565/119	based on extraction of label to an organic phase,
	i.e. partitioning of label between different organic
2565/125	phases
2565/125	. Electrophoretic separation
2565/131	Single/double strand conformational analysis, i.e. SSCP/DSCP
2565/133	conformational analysis
2565/137	Chromatographic separation
2565/20	Detection means characterised by being a gene
23 03/20	reporter based analysis
2565/201	Two hybrid system
2565/207	Three hybrid system
2565/30	Detection characterised by liberation or release of
	label
2565/301	Pyrophosphate (PPi)
2565/40	• Detection characterised by signal amplification of
	label
2565/401	Signal amplification by chemical polymerisation
2565/50	Detection characterised by immobilisation to a
2575/501	surface
2565/501	being an array of oligonucleotides
2565/507	characterised by the density of the capture oligonucleotide
2565/513	characterised by the pattern of the arrayed
20 00,010	oligonucleotides
2565/514	characterised by the use of the arrayed
	oligonucleotides as identifier tags, e.g. universal
	addressable array, anti-tag or tag complement
	array
2565/515	characterised by the interaction between or sequential use of two or more arrays
2565/518	characterised by the immobilisation of the nucleic
2505/510	acid sample or target
2565/519	characterised by the capture moiety being a single
	stranded oligonucleotide
2565/525	characterised by the capture oligonucleotide being
	double stranded
2565/531	characterised by the capture moiety being a protein for target oligonucleotides
2565/537	characterised by the capture oligonucleotide
2505/557	acting as a primer
2565/543	characterised by the use of two or more capture
	oligonucleotide primers in concert, e.g. bridge
	amplification
2565/549	characterised by the capture oligonucleotide being
0565160	a reporter labelled capture oligonucleotide
2565/60	Detection means characterised by use of a special device
2565/601	being a microscope, e.g. atomic force microscopy
2303/001	[AFM]
2565/607	• being a sensor, e.g. electrode
2565/619	being a video camera
2565/625	being a nucleic acid test strip device, e.g.
	dipsticks, strips, tapes, CD plates
2565/626	being a flow cytometer
2565/627	being a mass spectrometer
2565/628	being a surface plasmon resonance spectrometer
2565/629	being a microfluidic device
2565/631	being a biochannel or pore

2565/632	base a constant and a macanana Daman
	being a surface enhanced, e.g. resonance, Raman spectrometer
2565/633	NMR
2565/634	being an acoustic wave sensor
2600/00	Oligonucleotides characterized by their use
2600/106	• Pharmacogenomics, i.e. genetic variability in
	individual responses to drugs and drug metabolism
2600/112	 Disease subtyping, staging or classification
2600/118	 Prognosis of disease development
2600/124	 Animal traits, i.e. production traits, including
	athletic performance or the like
2600/13	• Plant traits
2600/136	Screening for pharmacological compounds
2600/142	. Toxicological screening, e.g. expression profiles
	which identify toxicity
2600/148	Screening for cosmetic compounds
2600/154	Methylation markers
2600/156	Polymorphic or mutational markers
2600/158	Expression markers
2600/16	 Primer sets for multiplex assays
2600/166	• Oligonucleotides used as internal standards, controls
	or normalisation probes
2600/172	. Haplotypes
2600/178	• miRNA, siRNA or ncRNA

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