

Product Description

SALSA® MLPA® Probemix P244 AIP-MEN1-CDKN1B

To be used with the MLPA General Protocol.

Version D1

For complete product history see page 10.

Catalogue numbers

- P244-025R: SALSA MLPA Probemix P244 AIP-MEN1-CDKN1B, 25 reactions.
- P244-050R: SALSA MLPA Probemix P244 AIP-MEN1-CDKN1B, 50 reactions.
- P244-100R: SALSA MLPA Probemix P244 AIP-MEN1-CDKN1B, 100 reactions.

SALSA® MLPA® Probemix P244 AIP-MEN1-CDKN1B (hereafter: P244 AIP-MEN1-CDKN1B) is to be used in combination with:

- 1. SALSA® MLPA® Reagent Kit (Cat. No: EK1-FAM, EK1-CY5, EK5-FAM, EK5-CY5, EK20-FAM),
- 2. Data analysis software Coffalyser.Net™ (Cat. No: n.a.)

Volumes and ingredients

Volumes			Ingradiente	
P244-025R	P244-050R	P244-100R	- Ingredients	
40 μΙ	80 µl	160 μΙ	Synthetic oligonucleotides, oligonucleotides purified from bacteria, Tris-HCl, EDTA	

The MLPA probemix is not known to contain any harmful agents. Based on the concentrations present, none of the ingredients are hazardous as defined by the Hazard Communication Standard. A Safety Data Sheet (SDS) is not required for this product: none of the ingredients contain dangerous substances at concentrations requiring distribution of an SDS (as per Regulation (EC) No 1272/2008 [EU-GHS/CLP] and 1907/2006 [REACH] and amendments).

Storage and handling

Recommended storage conditions	-25°C	*

A shelf life of until the expiry date is guaranteed, when stored in the original packaging under recommended conditions. For the exact expiry date, see the label on the vial. This product should not be exposed to more than 25 freeze-thaw cycles. Do not use the product if the packaging is damaged or opened. Leave chemicals in original containers. Waste material must be disposed of in accordance with the national and local regulations.

Certificate of Analysis

Information regarding quality tests and a sample electropherogram from the current sales lot is available at www.mrcholland.com.

Precautions and warnings

For professional use only. Always consult the most recent product description AND the MLPA General Protocol before use: www.mrcholland.com. It is the responsibility of the user to be aware of the latest scientific knowledge of the application before drawing any conclusions from findings generated with this product.

General information

SALSA® MLPA® Probemix P244-AIP-MEN1-CDKN1B is a **research use only (RUO)** assay for the detection of deletions or duplications in the *AIP*, *MEN1* and *CDKN1B* genes which are associated with familial isolated pituitary adenoma (FIPA), multiple endocrine neoplasia type 1 (MEN1) or multiple endocrine neoplasia type 4 (MEN4), respectively.

Multiple endocrine neoplasia type 1 (MEN1) is predominantly characterized by the occurrence of primary hyperparathyroidism (PHPT), which occurs in 95-100% of patients; pancreatic neuroendocrine tumours, which occur in 40-75% of patients; and pituitary adenoma, which is found in 30-50% of patients. Most tumours are non-metastasizing, but many can cause striking and serious clinical effects due to the increased secretion of hormones. It is estimated that in the general population 1 to 10 in 100.000 individuals develop MEN1 during their lifetime. Nine out of ten patients diagnosed with MEN1 have the familial form. MEN1 shows dominant autosomal inheritance and the penetrance is >95% by age 40 for confirmed pathogenic mutations. The mean age of death of MEN1 patients is between 50 and 55 years. The single gene associated with MEN1 syndrome is *MEN1*, which encodes the menin protein. Heterozygous *MEN1* pathogenic variants are found in ~90% of familial MEN1 syndrome patients and in ~65% of sporadic cases. Loss of heterozygosity (LOH) of MEN1 is observed in >90% MEN1 tumours suggesting that *MEN1* acts as a tumour suppressor gene, in line with the Knudson 2-hit hypothesis for tumorigenesis. Besides point mutations, several deletions involving one or more complete exons in the *MEN1* gene have been described (Carroll 2013, Concolino et al. 2016, Lemos and Thakker 2008, Romanet et al. 2019, Thakker 2014), including a pathogenic deletion of just the 5'-UTR (Kooblall et al. 2020).

Pituitary adenomas (PAs) occur with a frequency of ~1 in 1000 in the general population. Most cases are sporadic, but approximately 5% occurs as a familial cancer. The *AIP* gene encodes aryl hydrocarbon receptor-interacting protein (AIP), a tumour suppressor that is involved in the control of cell proliferation and differentiation. *AIP* loss of function mutations are found in 15-25% of familial isolated pituitary adenoma (FIPA) cases, which are subsequently referred to as *AIP*-FIPA. Inheritance is autosomal dominant and the average penetrance is 15-30%, although this may vary greatly. The prevalence of *AIP*-FIPA is estimated at 1:100,000. Similar as for *MEN1*, LOH is frequently observed, suggesting that *AIP* also acts as a tumour suppressor gene (Cai et al. 2013). Although most known germline *AIP* mutations are point mutations, several exon deletions have been reported: exon 1-2, exon 2, exon 1-6 (Georgitsi et al. 2008, Igreja et al. 2010, Marques et al. 2018).

MEN1 and AIP are located in close proximity on 11q13, and somatic LOH in MEN1 and FIPA associated tumours often affects both genes. Apart from tumours in MEN1 and FIPA patients, LOH of this locus also occurs in sporadic cancers, especially in endocrine tissues. As both genes are considered tumour suppressor genes this double loss may contribute to tumorigenesis. Chromosomal losses of the 11q13 chromosomal band have also been found in other cancers, such as cervical cancer and hibernomas (Newsham 1998; Nord et al. 2010).

MEN4 is a distinct MEN type but the symptoms of MEN4 largely overlap with MEN1 (Pellegata et al. 2006). In a small number (estimated at 1-3%) of *MEN1* mutation-negative patients fulfilling the diagnostic criteria for MEN1, mutations in *CDKN1B* have been detected. Extrapolating from this, the prevalence of MEN4 is very low: <1:300,000. Like MEN1, MEN4 is primarily characterized by PHPT and PA, but the additional tumours show some differences; tumours in the reproductive organs, and adrenal and renal tumours have been found in MEN4 patients. The only way to distinguish MEN4 from MEN1 is by identification of a pathogenic mutation in *CDKN1B*. Somatic mutations in *CDKN1B* have also been identified in sporadic tumours, but LOH of *CDKN1B* in MEN4-related tumours has not been found.

More information on MEN1 can be found on https://www.ncbi.nlm.nih.gov/books/NBK1538/

More information on AIP-related FIPA can be found on https://www.ncbi.nlm.nih.gov/books/NBK97965/.

More information on MEN4 can be found on: https://omim.org/entry/610755

This product is not CE/FDA registered for use in diagnostic procedures. The SALSA® MLPA® technique is covered by US patent 6,955,901 and corresponding patents outside the US. The purchase of this product includes a license to use only this amount of product solely for the purchaser's own use.



Gene structure and transcript variants:

Entrez Gene shows transcript variants of each gene: https://www.ncbi.nlm.nih.gov/gene
For NM_ mRNA reference sequences: https://www.ncbi.nlm.nih.gov/nuccore?db=nucleotide
Matched Annotation from NCBI and EMBL-EBI (MANE): https://www.ncbi.nlm.nih.gov/refseq/MANE

Exon numbering

The *MEN1* exon numbering used in this P244-AIP-MEN1-CDKN1B product description is the exon numbering derived from MANE project based on MANE Select transcript NM_001370259.2. The *AIP* exon numbering used is the exon numbering derived from MANE project based on MANE Select transcript NM_003977.4. The *CDKN1B* exon numbering used is the exon numbering derived from MANE project based on MANE Select transcript NM_004064.5. As changes to the databases can occur after release of this product description, the NM_ sequence and exon numbering may not be up-to-date.

Probemix content

P244-D1 AIP-MEN1-CDKN1B contains 42 MLPA probes with amplification products between 129 and 463 nucleotides (nt). This includes 25 probes for the *MEN1-AIP* region and five probes for the *CDKN1B* region. In addition, 12 reference probes are included that target relatively copy number stable regions in various cancer types. Complete probe sequences and the identity of the genes detected by the reference probes are available online (www.mrcholland.com).

This probemix contains nine quality control fragments generating amplification products between 64 and 105 nt: four DNA Quantity fragments (Q-fragments), two DNA Denaturation fragments (D-fragments), one Benchmark fragment, and one chromosome X and one chromosome Y-specific fragment (see table below). More information on how to interpret observations on these control fragments can be found in the MLPA General Protocol and online at www.mrcholland.com.

Length (nt)	Name
64-70-76-82	Q-fragments (only visible with <100 ng sample DNA)
88-96	D-fragments (low signal indicates incomplete denaturation)
92	Benchmark fragment
100	X-fragment (X chromosome specific)
105	Y-fragment (Y chromosome specific)

MLPA technique

The principles of MLPA (Schouten et al. 2002) are described in the MLPA General Protocol (www.mrcholland.com). More information on the use of MLPA in tumour applications can be found in Hömig-Hölzel and Savola (2012).

MLPA technique validation

Internal validation using 16 different DNA samples from healthy individuals is required, in particular when using MLPA for the first time, or when changing the sample type or the sample handling procedure, DNA extraction method or instruments used. This validation experiment should result in a standard deviation \leq 0.10 for all probes over the experiment.

Required specimens

Extracted DNA from human peripheral blood or DNA derived from fresh or fresh formalin-fixed, paraffinembedded (FFPE) tumour tissue, free from impurities known to affect MLPA reactions. MRC Holland has tested and can recommend the following extraction methods:

- QIAGEN Autopure LS (automated) and QIAamp DNA mini/midi/maxi kit (manual)
- Promega Wizard Genomic DNA Purification Kit (manual)
- Salting out (manual)

All samples tested, including reference DNA samples, should be derived from the same tissue type, handled using the same procedure, and prepared using the same DNA extraction method when possible. For more



information please refer to the section on DNA sample treatment found in the MLPA General Protocol. More information on the use of FFPE tissue samples for MLPA can be found in Atanesyan et al. (2017).

Reference samples

A sufficient number (≥3) of different reference samples from unrelated individuals should be included in each MLPA experiment for data normalisation. Reference samples should be derived from healthy individuals without a history of MEN or FIPA. More information regarding the selection and use of reference samples can be found in the MLPA General Protocol (www.mrcholland.com).

Positive control DNA samples

MRC Holland cannot provide positive DNA samples. Inclusion of a positive sample in each experiment is recommended. Coriell Institute (https://catalog.coriell.org) and Leibniz Institute DSMZ (https://www.dsmz.de) have diverse collections of biological resources which may be used as positive control DNA samples in your MLPA experiments. Sample ID number NA07981 from the Coriell Institute has been tested with this P244-D1 AIP-MEN1-CDKN1B at MRC Holland and can be used as a positive control sample to detect four copies of *CDKN1B* and the flanking regions. The quality of cell lines can change; therefore deviations to the indicated copy number alteration (CNA) findings might occur.

Data analysis

Coffalyser.Net should be used for data analysis in combination with the appropriate lot-specific Coffalyser sheet. For both, the latest version should be used. Coffalyser.Net is freely downloadable at www.mrcholland.com. Use of other non-proprietary software may lead to inconclusive or false results. For more details on MLPA quality control and data analysis, including normalisation, see the Coffalyser.Net Reference Manual.

Interpretation of results

The expected results for *MEN1*, *AIP* and *CDKN1B* target probes are allele copy numbers of 2 (normal), 1 (heterozygous deletion), 3 (heterozygous duplication), 0 (homozygous deletion) or 4 (homozygous duplication or heterozygous triplication). The standard deviation of each individual probe over all the reference samples should be ≤0.10 and the final ratio (FR) of each individual reference probe in the patient samples should be between 0.80 and 1.20. When these criteria are fulfilled, the following cut-off values for the FR of the probes can be used to interpret MLPA results for autosomal chromosomes or pseudo-autosomal regions:

Copy number status	Final ratio (FR)
Normal	0.80 < FR < 1.20
Homozygous deletion	FR = 0
Heterozygous deletion	0.40 < FR < 0.65
Heterozygous duplication/gain	1.30 < FR < 1.65
Heterozygous triplication/homozygous duplication/gain	1.75 < FR < 2.15
Ambiguous copy number	All other values

Note: The term "dosage quotient", used in older product description versions, has been replaced by "final ratio" to become consistent with the terminology of Coffalyser.Net (Calculations, cut-offs and interpretation remain unchanged.) Please note that Coffalyser.Net also shows arbitrary borders as part of the statistical analysis of results obtained in an experiment. As such, arbitrary borders are different from the final ratio cut-off values shown here above.

Please note that these above mentioned final ratios are only valid for germline testing. Final ratios are affected both by percentage of tumour cells and by possible subclonality.

- <u>Arranging probes</u> according to chromosomal location facilitates interpretation of the results and may reveal more subtle changes such as those observed in mosaic or subclonal cases. Analysis of parental samples may be necessary for correct interpretation of complex results.
- <u>False positive results</u>: Please note that abnormalities detected by a single probe (or multiple consecutive probes) still have a considerable chance of being a false positive result. Sequence changes (e.g. single

nucleotide variants, point mutations) in the target sequence detected by a probe can be one cause. Incomplete DNA denaturation (e.g. due to salt contamination in the DNA sample) can also lead to a decreased probe signal, in particular for probes located in or near a GC-rich region or in or near the *MEN1-AIP* gene. The use of an additional purification step or an alternative DNA extraction method may resolve such cases. Additionally, contamination of DNA samples with cDNA or PCR amplicons of individual exons can lead to an increased probe signal (Varga et al. 2012). Analysis of an independently collected secondary DNA sample can exclude these kinds of contamination artefacts.

- Normal copy number variation in healthy individuals is described in the database of genomic variants: http://dgv.tcag.ca/dgv/app/home. Users should always consult the latest update of the database and scientific literature when interpreting their findings.
- Not all abnormalities detected by MLPA are pathogenic. In some genes, intragenic deletions are known that result in very mild or no disease (as described for DMD by Schwartz et al. 2007). For many genes, more than one transcript variant exists. Copy number changes of exons that are not present in all transcript variants may not have clinical significance. Duplications that include the first or last exon of a gene (e.g. exons 1-3) might not result in inactivation of that gene copy.
- Copy number changes detected by reference probes or flanking probes are unlikely to have any relation to the condition tested for.
- False results can be obtained if one or more peaks are off-scale. For example, a duplication of one or more exons can be obscured when peaks are off-scale, resulting in a false negative result. The risk on off-scale peaks is higher when probemixes are used that contain a relatively low number of probes. Coffalyser.Net software warns for off-scale peaks while other software does not. If one or more peaks are off-scale, rerun the PCR products using either: a lower injection voltage or a shorter injection time, or a reduced amount of sample by diluting PCR products.

Limitations of the procedure

- In most populations, the major cause of genetic defects in the *MEN1*, *AIP* and *CDKN1B* gene are small (point) mutations, none of which will be detected by using P244 AIP-MEN1-CDKN1B.
- MLPA cannot detect any changes that lie outside the target sequence of the probes and will not detect copy number neutral inversions or translocations. Even when MLPA did not detect any aberrations, the possibility remains that biological changes in that gene or chromosomal region *do* exist but remain undetected.
- Sequence changes (e.g. SNVs, point mutations) in the target sequence detected by a probe can cause false positive results. Mutations/SNVs (even when >20 nt from the probe ligation site) can reduce the probe signal by preventing ligation of the probe oligonucleotides or by destabilising the binding of a probe oligonucleotide to the sample DNA.
- MLPA analysis on tumour samples provides information on the average situation in the cells from which the DNA sample was purified. Gains or losses of genomic regions or genes may not be detected if the percentage of tumour cells is low. In addition, subclonality of the aberration affects the final ratio of the corresponding probe. Furthermore, there is always a possibility that one or more reference probes do show a copy number alteration in a patient sample, especially in tumours with high chromosomal instability.

Confirmation of results

Copy number changes detected by only a single probe always require confirmation by another method. An apparent deletion detected by a single probe can be due to e.g. a mutation/polymorphism that prevents ligation or destabilises the binding of probe oligonucleotides to the DNA sample. Sequence analysis can establish whether mutations or polymorphisms are present in the probe target sequence. The finding of a heterozygous mutation or polymorphism in sequence data indicates that two different alleles of the sequence are present in the sample DNA and that a false positive MLPA result was obtained.

Copy number changes detected by more than one consecutive probe should be confirmed by another independent technique such as long range PCR, qPCR, array CGH or Southern blotting, whenever possible. Deletions/duplications of more than 50 kb in length can often be confirmed by FISH.



Mutation databases

The UMD-MEN1 mutation database (www.umd.be/MEN1/) and the LOVD page for *MEN1*: databases.lovd.nl/shared/genes/MEN1. We strongly encourage users to deposit positive results in any of the *MEN1* databases. Recommendations for the nomenclature to describe deletions/duplications of one or more exons can be found on http://varnomen.hgvs.org/.

The FIPA Patients *AIP* Mutations Database: https://aip.fipapatients.org/menu/main/home and the LOVD page for *AIP*: https://databases.lovd.nl/shared/genes/AIP. We strongly encourage users to deposit positive results in any of the *AIP* mutation databases. Recommendations for the nomenclature to describe deletions/duplications of one or more exons can be found on http://varnomen.hgvs.org/.

The LOVD page for *CDKN1B*: https://databases.lovd.nl/shared/genes/CDKN1B. We strongly encourage users to deposit positive results in the *CDKN1B* database. Recommendations for the nomenclature to describe deletions/duplications of one or more exons can be found on http://varnomen.hgvs.org/.

COSMIC mutation database

Somatic mutations in *MEN1*, *AIP* and *CDKN1B* in cancer can be found in the COSMIC database at https://cancer.sanger.ac.uk/cosmic.

Please report copy number changes detected by the reference probes, false positive results due to SNVs and unusual results (e.g., a duplication of *AIP* exons 3 and 5 but not exon 4 to MRC Holland: info@mrcholland.com.



Table 1. P244-D1 AIP-MEN1-CDKN1B

onath (nt)	CALCA MI DA probo	Chro	Chromosomal position (hg18) ^a		
ength (nt)	SALSA MLPA probe	Reference	MEN1-AIP region	CDKN1B	
64-105	Control fragments – see table in prob	emix content section	for more information		
129 *	Reference probe 18709-L26847	5q31			
136 ¥ «	MEN1 probe 22342-L02795		Exon 9		
142 * ¬	GPRC5A probe 22344-L03556			Downstream	
148	CDKN1B probe 18425-L23733			Exon 2	
154 *	Reference probe 04540-L28278	2q24			
160 « ¬	BRMS1 probe 04155-L03510		11q13		
167 «	MEN1 probe 13158-L14680		Exon 4		
175 «	AIP probe 07380-L09559		Exon 1		
184 *	Reference probe 10904-L27810	9q34			
191 «	MEN1 probe 01663-L01242		Upstream		
195 «	MEN1 probe 18765-L24187		Exon 6		
202 «	MEN1 probe 13159-L14681		Exon 5		
209 «	AIP probe 07383-L07030		Exon 4		
216 *	Reference probe 09103-L31536	4q25			
220 «	MEN1 probe 01664-L01243		Exon 2		
229 « ¬	RELA probe 01120-L00060		11q13		
238 «	AIP probe 07381-L07028		Exon 2		
247 «	MEN1 probe 01164-L00720		Exon 10		
256 *	Reference probe 08812-L08872	2p13			
266 «	AIP probe 07384-L09556		Exon 5		
277 *	Reference probe 13393-L28135	6q12			
283 «	MEN1 probe 01665-L14816		Exon 3		
292 «	AIP probe 07379-L09558		Exon 1		
301 «	MEN1 probe 01666-L01245		Exon 7		
310 ¬	CCND1 probe 05403-L04809		11q13		
319 «	AIP probe 07385-L09557		Exon 6		
326 *	Reference probe 16275-L18567	19p13			
338 ¥ ¬	SNX15 probe 01667-L31522		11q13		
346 *	Reference probe 04337-L20895	15q21			
355 « ¬	FAM89B probe 04157-L03512		11q13		
364 «	AIP probe 07382-L09069		Exon 3		
373 *	Reference probe 04278-L03682	12q12			
382 ¬	SART1 probe 04159-L03514		11q13		
391	CDKN1B probe 18426-L23497			Exon 1	
400 « +	MEN1 probe 18427-L23498		Upstream		
409 *	Reference probe 14839-L30627	1p34			
420 * « ¬	SF1 probe 22341-L20815		11q13		
427 «	CDKN1B probe 18429-L23500		-	Exon 3	
436 «	MEN1 probe 18430-L23501		Exon 8		
445 *	Reference probe 10709-L11291	6p12			
454 * ¬	BCL2L14 probe 22345-L02547			Upstream	
463 *	Reference probe 15970-L18122	18p11			

^a See section Exon numbering on page 2 for more information.

^{*} New in version D1.

[¥] Changed in version D1. Minor alteration, no change in sequence detected.

[«] Probe located in or near a GC-rich region. A low signal can be caused by salt contamination in the DNA sample leading to incomplete DNA denaturation, especially of GC-rich regions.

[¬] Flanking probe. Included to help determine the extent of a deletion/duplication. Copy number alterations of only the flanking or reference probes are unlikely to be related to the condition tested.

⁺ The ligation site of this probe is located in an alternative exon 1 present in two alternative transcripts: NM_130803.2 and NM_130804.2. In both transcripts the ligation site is on position 242-243. The significance of deletions/duplications of





only this alternative exon is not clear as this exon is non-coding and alternative transcript variants using other transcription start sites are known.

The probe lengths in the table above may vary slightly depending on the capillary electrophoresis machine settings. Please see the most up-to-date Coffalyser sheet for exact probe lengths obtained at MRC Holland.

SNVs located in the target sequence of a probe can influence probe hybridization and/or probe ligation. Single probe aberration(s) must be confirmed by another method.

Table 2. Target and flanking probes arranged according to chromosomal location

Table 2a. MEN1-AIP region (11q13)

Length (nt)	SALSA MLPA probe	Gene exon ^a	Ligation site	Partial sequence ^b (24 nt adjacent to ligation site)	Distance to next probe
420 « ¬	22341-L20815	SF1		GATTCCAGGAAT-GCCTACAGTTAT	27.9 kb
		MEN1	NM_001370259.2		
		stop codon	1892-1894 (exon 10)		
247 « »	01164-L00720	Exon 10	1791-1792	CGCCATCAAGCT-GCAACTCACGGC	0.7 kb
136 « »	22342-L02795	Exon 9	1326-1327	CGGCATCTGCAA-ATGGGAGGAGGG	0.6 kb
436 «	18430-L23501	Exon 8	1164-1165	CTTTGAAGTAGC-CAATGATGTCAT	0.6 kb
301 « »	01666-L01245	Exon 7	1028-1029	ACCCCTACATGT-ACCTGGCTGGCT	0.7 kb
195 « »	18765-L24187	Exon 6	972-973	CCTCTACCACAA-GGTGGGGGCATC	0.2 kb
202 « »	13159-L14681	Exon 5	867-868	GCTGCTCTATGA-CCTGGGACATCT	0.4 kb
167 « »	13158-L14680	Exon 4	761-762	GTGACCGCAAGA-TGGAGGTGGCGT	0.4 kb
283 « »	01665-L14816	Exon 3	608-609	AGGATCATGCCT-GGGTAGTGTTTG	2.0 kb
220 « »	01664-L01243	Exon 2	201-202	CGTGGAGCATTT-TCTGGCTGTCAA	0.7 kb
		start codon	62-64 (exon 2)		
191 « »	01663-L01242	Upstream	131 nt before exon 1	GAGATCCCAGAA-GCCACAGCGCAG	0.4 kb
400 « +	18427-L23498	Upstream	501 nt before exon 1	GCGGAAGTGGGA-AACGAGTGCTGC	216.5 kb
338 ¬	01667-L31522	SNX15		CGAAGGATGACT-TCCTGCGGCACT	545.9 kb
355 « ¬	04157-L03512	FAM89B		ACAAACACCTGT-GCCAAGACCTGA	88.3 kb
229 « ¬	01120-L00060	RELA		AAAGGACTGCCG-GGATGGCTTCTA	305.6 kb
382 ¬	04159-L03514	SART1		CCGCAAGAAGGA-GAAGGAGGTAGT	374.9 kb
160 « ¬	04155-L03510	BRMS1		CAGAAGAGATGG-AAGCAGAGGGTG	1141.0 kb
		AIP	NM_003977.4		
292 «	07379-L09558	Exon 1	105-106	GAGTCCGGAAGT-TGCCGAAAGGGA	0.1 kb
		start codon	134-136 (exon 1)		
175 «	07380-L09559	Exon 1	186-187	AAAACGTGTGAT-ACAGGAAGGCCG	3.9 kb
238 «	07381-L07028	Exon 2	320-321	CCATGGAGCTCA-TCATTGGCAAGA	2.3 kb
364 «	07382-L09069	Exon 3	519-520	ACAGATGCGTGA-ACACAGCTCCCT	0.7 kb
209 «	07383-L07030	Exon 4	676-677	GCAGTGCCACTT-ATCCACCAGGAG	0.3 kb
266 «	07384-L09556	Exon 5	840-841	GCTGCTGCTCAA-CTACTGCCAGTG	0.5 kb
319 «	07385-L09557	Exon 6	997-998	CAGGCTGACTTT-GCCAAAGTGCTG	2157.1 kb
		stop codon	1124-1126 (exon 6)		
310 ¬	05403-L04809	CCND1		TCCGCCCTCCAT-GGTGGCAGCGGG	-

Table 2b. CDKNB1

Length (nt)	SALSA MLPA probe	CDKN1B exon ^a	Ligation site NM_004064.5	Partial sequence ^b (24 nt adjacent to ligation site)	Distance to next probe
454 ¬	22345-L02547	BCL2L14		GTCATGGACAGA-GGTGTCATGGCC	638.3 kb
391	18426-L23497	Exon 1	470-471	GACCCGGGAGAA-AGATGTCAAACG	1.1 kb
		start codon	473-475 (exon 1)		
148	18425-L23733	Exon 2	1022-1023	ATGCCGGTTCTG-TGGAGCAGACGC	2.3 kb
		stop codon	1067-1069 (exon 2)		
427 «	18429-L23500	Exon 3	1199-1200	CCTGTATAAGCA-CTGAAAAACAAC	187.3 kb
142 ¬	22344-L03556	GPRC5A		GTCTGCAAGGTG-CAGGACTCCAAC	-

^a See section Exon numbering on page 2 for more information.

- « Probe located in or near a GC-rich region. A low signal can be caused by salt contamination in the DNA sample leading to incomplete DNA denaturation, especially of GC-rich regions.
- » Detects the same sequence as MEN1 probes in SALSA MLPA Probemix P017-D1 MEN1.
- ¬ Flanking probe. Included to help determine the extent of a deletion/duplication. Copy number alterations of only the flanking or reference probes are unlikely to be related to the condition tested.
- + The ligation site of this probe is located in an alternative exon 1 present in two alternative transcripts: NM_130803.2 and NM_130804.2. In both transcripts the ligation site is on position 242-243. The significance of deletions/duplications of only this alternative exon is not clear as this exon is non-coding and alternative transcript variants using other transcription start sites are known.

SNVs located in the target sequence of a probe can influence probe hybridization and/or probe ligation. Single probe aberration(s) must be confirmed by another method.

Related products

For related products, see the product page on our website.

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^b Only partial probe sequences are shown. Complete probe sequences are available at www.mrcholland.com. Please notify us of any mistakes: info@mrcholland.com.



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Selected publications using P244 AIP-MEN1-CDKN1B

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- Carvalho RA et al. (2018). Germline mutation landscape of multiple endocrine neoplasia type 1 using full gene next-generation sequencing. *Eur J Endocrinol*. 179:391-407.
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- Romanet P et al. (2019). UMD-MEN1 Database: An Overview of the 370 MEN1 Variants Present in 1676 Patients From the French Population. *J Clin Endocrinol Metab*. 104:753-764.
- Tuncer FN et al. (2018). Screening of AIP Gene Variations in a Cohort of Turkish Patients with Young-Onset Sporadic Hormone-Secreting Pituitary Adenomas. *Genet Test Mol Biomarkers*. 22:702-708.

MEN1 copy number variations have also been detected with the P017 MEN1 probemix. See the selected publications listed in the P017 product description for relevant publications.

P244 prod	P244 product history		
Version	Modification		
D1	Three new flanking probes have been included and the length of two probes was changed. All reference probes were replaced and the total number of reference probes was increased to twelve.		
C1	Three probes for the missing <i>MEN1</i> exons were included and probes for each exon of the <i>CDKN1B</i> gene are included. Seven reference probes were replaced and one extra reference probe was included.		
B1	One MEN1 probe was removed and two new MEN1 probes were included. Several reference probes were replaced and four extra control fragments at 88-96-100-105 nt were included.		
A1	First release.		





Implemented changes in the product description

Version D1-04 - 30 October 2025 (05P)

- Product description updated as P244 is for research use only from lot 0425 onwards.
- Product description adapted to a new template.

Version D1-03 - 22 April 2025 (04P)

- List of Selected Publications shortened.
- In Table 1, the MEN1-AIP region column has been updated based on MANE exon numbering. In Table 2a, the gene exon and ligation site columns have been updated based on MANE exon numbering. The *Transcript variants* and *Exon numbering* sections have also been edited to reflect this change.

Version D1-02 - 03 February 2021 (04P)

- Product description rewritten and adapted to a new template.
- Intended Purpose was rewritten.
- NM sequences for AIP and CDKN1B genes were updated and the ligation sites were also updated according to these new versions of the NM reference sequences in Table 2.
- References and Selected Publications were curated and new literature was included.
- UK has been added to the list of countries in Europe that accept the CE mark.

More information: www.mrcholland.com; www.mrcholland.eu		
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IVD	EUROPE* (until lot 0822)
RUO	EUROPE* (as of lot 0425) ALL OTHER COUNTRIES

^{*}comprising EU (candidate) member states and members of the European Free Trade Association (EFTA), and the UK. The product is for RUO in all other European countries.

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