

Product Description SALSA[®] MLPA[®] Probemix P106-D2 X-linked ID

To be used with the MLPA General Protocol.

Version D2

As compared to version D1, the length of one probe has been adjusted. For complete product history see page 13.

Catalogue numbers:

- P106-025R: SALSA MLPA Probemix P106 X-linked ID, 25 reactions.
- P106-050R: SALSA MLPA Probemix P106 X-linked ID, 50 reactions.
- P106-100R: SALSA MLPA Probemix P106 X-linked ID, 100 reactions.

To be used in combination with a SALSA MLPA reagent kit and Coffalyser.Net data analysis software. MLPA reagent kits are either provided with FAM or Cy5.0 dye-labelled PCR primer, suitable for Applied Biosystems and Beckman/SCIEX capillary sequencers, respectively (see www.mrcholland.com).

Certificate of Analysis

Information regarding storage conditions, 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

The SALSA MLPA Probemix P106 X-linked ID is a **research use only (RUO)** assay for the detection of deletions or duplications in several genes on the X-chromosome, which are associated with X-linked intellectual disability (XLID).

Intellectual disability (ID) is estimated to occur in about 2-3% of the population, and 12% of all ID is thought to be XLID (Utine et al. 2011). Among intellectually disabled patients, an excess of males over females has long been noted, which is usually explained by the presence of many genes responsible for ID on the X chromosome (Ropers et al. 2005).

XLID is usually divided into a syndromic and a non-syndromic form. In syndromic forms (S-XLID), ID is present in association with a specific pattern of physical, neurological, and/or metabolic abnormalities. The term nonspecific or non-syndromic XLID (NS-XLID) was introduced to indicate a condition segregating in an X-linked manner in which male patients have no consistent phenotypic manifestations other than ID. Many different genes responsible for both forms of XLID have been identified (Ropers et al. 2005).

This SALSA MLPA probemix is not CE/FDA registered for use in diagnostic procedures. Purchase of this product includes a limited license for research purposes.

Gene structure and transcript variants:

Entrez Gene shows transcript variants of each gene: http://www.ncbi.nlm.nih.gov/sites/entrez?db=gene For NM_ mRNA reference sequences: http://www.ncbi.nlm.nih.gov/sites/entrez?db=nucleotide Locus Reference Genomic (LRG) database: http://www.lrg-sequence.org/



Exon numbering

The exon numbering used in this P106-D2 X-linked ID product description for the different targeted genes is the exon numbering from the following NG and LRG sequences:

Gene name	NG / LRG sequence
RPS6KA3	NG_007488.1
ARX	NG_008281.1
IL1RAPL1	NG_008292.2
TSPAN7	NG_009160.1
PQBP1	NG_015967.1
HUWE1	NG_016261.2
OPHN1	NG_008960.1
ACSL4	NG_008053.1
PAK3	NG_008288.2
DCX	NG_011750.1
AGTR2	NG_016326.1
ARHGEF6	NG_008873.1
FMR1	LRG_762
AFF2	NG_016313.2
SLC6A8	NG_012016.2
GDI1	NG_008954.1

The exon numbering of the NM_ sequence that was used for determining a probe's ligation site does not always correspond to the exon numbering obtained from the NG or LRG sequences. 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

The SALSA MLPA Probemix P106-D2 X-linked ID contains 46 MLPA probes with amplification products between 130 and 481 nucleotides (nt). These probes detect sequences in 16 genes located in the X-chromosome. Complete probe sequences are available online (www.mrcholland.com).

This probemix contains ten quality control fragments generating amplification products between 64 and 121 nt: four DNA Quantity fragments (Q-fragments), two DNA Denaturation fragments (D-fragments), one Benchmark fragment, and one chromosome X and two chromosome Y-specific fragments (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)			
105-121	Y-fragments (Y chromosome specific)			



MLPA technique

The principles of the MLPA technique (Schouten et al. 2002) are described in the MLPA General Protocol (www.mrcholland.com).

MLPA technique validation

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

Required specimens

Extracted DNA, free from impurities known to affect MLPA reactions. For more information please refer to the section on DNA sample treatment found in the MLPA General Protocol.

Reference samples

A sufficient number (\geq 3) of reference samples should be included in each MLPA experiment for data normalisation. 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. Reference samples should be derived from different unrelated individuals who are from families without a history of XLID. It is recommended to use samples of the same sex to facilitate interpretation. 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. The quality of cell lines can change; therefore samples should be validated before use.

Data analysis

Coffalyser.Net software should be used for data analysis in combination with the appropriate lot-specific MLPA Coffalyser sheet. For both, the latest version should be used. Coffalyser.Net software 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 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:

Copy Number status: Male samples	Final ratio
Normal	0.80 < FR < 1.20
Deletion	FR = 0
Duplication	1.65 < FR < 2.25
Ambiguous copy number	All other values



Copy Number status: Female samples	Final ratio
Normal	0.80 < FR < 1.20
Homozygous deletion	FR = 0
Heterozygous deletion	0.40 < FR < 0.65
Heterozygous duplication	1.30 < FR < 1.65
Heterozygous triplication/homozygous duplication	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 the Coffalyser.Net software. (Calculations, cut-offs and interpretation remain unchanged.) Please note that the Coffalyser.Net software 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.

- <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 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. SNVs, point mutations) in the target sequence detected by a probe can be one cause. Incomplete DNA denaturation (e.g. due to salt contamination) 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 *SLC6A8*, *ARX*, *TSPAN7*, *AFF2* and *GDI1* genes. 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.
- <u>Normal copy number variation</u> in healthy individuals is described in the database of genomic variants: <u>http://dgv.tcag.ca/dgv/app/home</u>. 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.
- <u>False results can be obtained if one or more peaks are off-scale</u>. 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 most of the genes targeted by this P106 X-linked ID probemix are small (point) mutations, most of which will not be detected by using SALSA MLPA Probemix P106 X-linked ID.
- 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.

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 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.

Database of genomic variation and phenotype in humans using Ensembl resources (DECIPHER)

https://decipher.sanger.ac.uk/. We strongly encourage users to deposit positive results in the DECIPHER database. Recommendations for the nomenclature to describe deletions/duplications of one or more exons can be found on http://varnomen.hgvs.org/.

Please report copy number changes detected by the reference probes, false positive results due to SNVs and unusual results (e.g., a duplication of *IL1RAPL1* exons 1 and 3 but not exon 2) to MRC Holland: info@mrcholland.com.



Table 1. SALSA MLPA Probemix P106-D2 X-linked ID

Length (nt)	SALSA MLPA probe	Gene detected	Chromosomal position (hg18)	Location (hg18) in kb
64-121	Control fragments – see ta	ble in probemix conten	t section for more information	
130	13917-L02320	AGTR2	Xq23	X-115,218
137 «	16875-L19669	GDI1	Xq28	X-153,319
142	02928-L03720	FMR1	Xq27.3	X-146,835
148 «	02935-L02326	ACSL4	Xq22.3	X-108,798
154 «	03511-L04202	AFF2	Xq28	X-147,390
160 «	02903-L02297	TSPAN7	Xp11.4	X-038,306
166	02927-L03721	FMR1	Xq27.3	X-146,822
172	16857-L19651	ARHGEF6	Xq26.3	X-135,617
178	02907-L02301	RPS6KA3	Xp22.12	X-020,084
184	13919-L15456	HUWE1	Xp11.22	X-053,598
188 «	22782-L32127	SLC6A8	Xq28	X-152,614
195 «	18790-L24221	ARX	Xp21.3	X-024,941
202	02902-L04460	ARHGEF6	Xq26.3	X-135,585
208 «	01871-L15827	SLC6A8	Xq28	X-152,610
215	04123-L15828	DCX	Xq23	X-110,531
222 «	02898-L04200	ARX	Xp21.3	X-024,935
229 «	13669-L15822	ARX	Xp21.3	X-024,944
235	02922-L23556	IL1RAPL1	Xp21.3	X-029,211
241	03516-L15823	AFF2	Xq28	X-147,727
248 «	03512-L23557	ACSL4	Xq22.3	X-108,863
256	13920-L23672	HUWE1	Xp11.22	X-053,691
263	02933-L23673	AFF2	Xq28	X-147,877
268 ~	02904-L23558	TSPAN7	Xp11.4	X-038,420
275	04124-L03481	DCX	Xq22.3	X-110,463
283	00493-L00066	AFF2	Xq28	X-147,551
288 ¥	23008-L02314	IL1RAPL1	Xp21.3	X-028,516
301	22783-L32128	ACSL4	Xq22.3	X-108,774
309	22016-L02878	PQBP1	Xp11.23	X-048,641
319	04121-L08390	DCX	Xq23	X-110,541
328	02921-L02315	IL1RAPL1	Xp21.3	X-028,717
337	02932-L02323	AFF2	Xq28	X-147,845
343	22017-L02293	ARHGEF6	Xq26.3	X-135,691
355	02925-L02319	AGTR2	Xq23	X-115,216
364	02906-L02300	RPS6KA3	Xp22.12	X-020,137
371	02912-L02306	OPHN1	Xq12	X-067,570
378 «	16874-L23559	GDI1	Xq28	X-153,323
385	02908-L03178	PAK3	Xq22.3	X-110,253
393	22856-L32371	PQBP1	Xp11.23	X-048,644
400	03521-L02304	PAK3	Xq22.3	X-110,346
409	02913-L23560	OPHN1	Xq12	X-067,436
418	02909-L02303	PAK3	Xq22.3	X-110,293
427	02923-L23561	IL1RAPL1	Xp21.2	X-029,596
436	02914-L02308	OPHN1	Xq12	X-067,334
443	16856-L19650	ARHGEF6	Xq26.3	X-135,655
472	02915-L02309	OPHN1	Xq12	X-067,201
481	02911-L02305	PAK3	Xq22.3	X-110,350

 $\ensuremath{\texttt{\textbf{Y}}}$ Changed in version D2. 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.

~ More variable. This probe has been reported to be deleted/duplicated in healthy individuals (various reports).



start codon

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. P106-D2 probes arranged according to chromosomal location

Tuble 2							
Length (nt)	SALSA MLPA probe	RPS6KA3 exonª	Ligation site NM_004586.3	Partial sequence ^b (24 nt adjacent to ligation site)	Distance to next probe		
Some truncat disabil	Some mutations in the <i>RPS6KA3</i> gene, cause mild intellectual disability (MRX19). Most mutations (incl. truncating) cause Coffin-Lowry syndrome (CLS). CLS is characterised by (amongst others) intellectual disability, fleshy hands and prominent ears. For more information on Coffin-Lowry syndrome see https://www.ncbi.nlm.nih.gov/books/NBK1346/.						
	stop codon 2498-2500 (Exon 22)						
178	02907-L02301	Exon 21	2282-2283	ATCAGAGACTGA-CTGCTGCTCTTG	53.1 kb		
364	02906-L02300	Exon 3	457-458	AAGGAAGGACAT-GAAAAGGCAGAT	4.8 M b to ARX gene		

278-280 (Exon 1)

Table 2a. RPS6KA3 gene. Xp22.12

Table 2b. ARX gene, Xp21.3

Length (nt)	SALSA MLPA probe	ARX exon ^a	Ligation site NM_139058.3	Partial sequence ^b (24 nt adjacent to ligation site)	Distance to next probe	
syndro syndro Parting	Mutations in the <i>ARX</i> gene underlie a phenotypic spectrum and can cause X-linked infantile spasm syndrome (https://omim.org/entry/308350), lissencephaly (https://omim.org/entry/300215), Proud syndrome (https://omim.org/entry/300004), intellectual disability (https://omim.org/entry/300419) and Partington syndrome (https://omim.org/entry/309510).					
P189 C	DKL5/ARX/FOX	G1: contains	more probes for the A	RX gene.		
		stop codon	1915-1917 (Exon 5)			
222 «	02898-L04200	Exon 4	1652-1651 reverse	GGCTGATGAAAG-CTGGGTGTCGGA	5.9 kb	
195 «	18790-L24221	Exon 2	1168-1167 reverse	GCTGCCCGCAGA-GAGGCACACGCT	2.8 kb	
229 «	13669-L15822	Exon 1	115-116	AGATCGCAATAA-TATCCGTTATAA	3.6 M b to IL1RAPL1 gene	
		start codon	229-231 (Exon 1)			

Table 2c. IL1RAPL1 gene, Xp21.3 - p21.2

Length (nt)	SALSA MLPA probe	IL1RAPL1 exonª	Ligation site NM_014271.4	Partial sequence ^b (24 nt adjacent to ligation site)	Distance to next probe		
	Mutations and/or deletions in the <i>IL1RAPL1</i> gene have been identified in families with X-linked syndromic intellectual disability (MRX21, see https://www.omim.org/entry/300143).						
		start codon	627-629 (Exon 2)				
288	23008-L02314	Exon 1	363-364	CAGCAAACAATC-GGGCACTTTGAG	201.5 kb		
328	02921-L02315	Exon 2	636-637	AGATGAAAGCTC-CGATTCCACACT	493.7 kb		
235	02922-L23556	Exon 3	834-835	TTGCCCAAAGTG-CTGGACTCAGTT	385.4 kb		
427	02923-L23561	Exon 6	1344-1345	CTCTGACTGATA-AGCCACCCAAGC	8.7 M b to <i>TSPAN7</i> gene		
		stop codon	2715-2717 (Exon 11)				

Table 2d. TSPAN7 gene, Xp11.4

Length (nt)	SALSA MLPA probe	TSPAN7 exonª	Ligation site NM_004615.4	Partial sequence ^b (24 nt adjacent to ligation site)	Distance to next probe		
	(Truncating) mutations in the <i>TSPAN7</i> gene have been identified as the cause of intellectual disability (MRX58, see https://www.omim.org/entry/300210).						
		start codon	6-8 (Exon 1)				
160 «	02903-L02297	Exon 1	32-33	ATGGAGACCAAA-CCTGTGATAACC	114.1 kb		

Length (nt)	SALSA MLPA probe	TSPAN7 exonª	Ligation site NM_004615.4	Partial sequence ^b (24 nt adjacent to ligation site)	Distance to next probe
268 ~	02904-L23558	Exon 5	467-468	TGTGGTGTGCAG-AACTACACCAAC	10.2 M b to <i>PQBP1</i> gene
		stop codon	753-755 (Exon 7)		

Table 2e. PQBP1 gene, Xp11.23

Length (nt)	SALSA MLPA probe	PQBP1 exonª	Ligation site NM_005710.2	Partial sequence ^b (24 nt adjacent to ligation site)	Distance to next probe	
	Mutations in the PQBP1 gene have been identified as the cause of Renpenning s (https://www.omim.org/entry/309500).					
		start codon	255-257 (Exon 1)			
309	22016-L02878	Exon 1	155-156	AGATGAGTACAT-GTTTACGGGAGG	3.8 kb	
393	22856-L32371	Exon 4	567-568	AAAAGTTGGACC-GGAGCCATGACA	5.0 M b to HUWE1 gene	
		stop codon	1050-1052 (Exon 6)			

Table 2f. HUWE1 gene, Xp11.22

Length (nt)	SALSA MLPA probe	HUWE1 exonª	Ligation site NM_031407.7	Partial sequence ^b (24 nt adjacent to ligation site)	Distance to next probe		
intelle intelle	Mutations in the <i>HUWE1</i> gene have been identified as the cause of Turner type of X-linked syndromic intellectual disability (MRXST, see https://omim.org/entry/309590). A nonsyndromic form of X-linked intellectual disability (MRX17 or MRX31, see https://www.omim.org/entry/309590). A nonsyndromic form of X-linked intellectual disability (MRX17 or MRX31, see https://www.omim.org/entry/309590). A nonsyndromic form of X-linked intellectual disability (MRX17 or MRX31, see https://www.omim.org/entry/300705) is caused by microduplications of chromosome Xp11.22, which includes the <i>HUWE1</i> gene.						
		stop codon	13516-13518 (Exon 84)				
184	13919-L15456	Exon 61	8669-8670	ATCTGAGTCCAA-GGAGACCCTTGG	92.6 kb		
256	13920-L23672	Exon 6	600-601	GCAGATGCTGGA-CAGACAGTGGAG	13.5 M b to OPHN1 gene		
		start codon	394-396 (Exon 4)				

Table 2g. OPHN1 gene, Xq12

Length (nt)	SALSA MLPA probe	OPHN1 exonª	Ligation site NM_002547.3	Partial sequence ^b (24 nt adjacent to ligation site)	Distance to next probe
Mutations/deletions in the <i>OPHN1</i> gene cause X-linked intellectual disability with distinctive appearance and cerebellar hypoplasia (https://omim.org/entry/300486).					
аррса		stop codon	2739-2741 (Exon 24)		
472	02915-L02309	Exon 21	2209-2210		
436	02914-L02308	Exon 12	1389-1390	AGGCCCTTTCAG-AAGCTAACAGAA	101.8 kb
409 #	02913-L23560	Exon 3	527-528	CAGACGCTGCAG-TCATTTCAGTTT	134.3 kb
371	02912-L02306	Exon 1	173-174	TGCTGCTTATCT-GGGAAGGCGATG	41.2 Mb to ACSL4 gene
		start codon	333-335 (Exon 2)		

Table 2h. ACSL4 gene, Xq22.3

Length (nt)	SALSA MLPA probe	ACSL4 exon ^a	Ligation site NM_022977.3	Partial sequence ^b (24 nt adjacent to ligation site)	Distance to next probe
It has been suggested that mutations in the ACSL4 gene might play a role in the development of inte disability (MRX63, see https://www.omim.org/entry/300387).					
		stop codon	2504-2506 (Exon 17)	2504-2506 (Exon 17)	
301	22783-L32128	Exon 17	2473-2474	AACCATTACCTC-AAAGACATTGAA	24.1 kb
148 «	02935-L02326	Exon 12	1745-1746	ATGTCTGCTTCT-GCTGCCCAATTG	65.1 kb
248 «	03512-L23557	Exon 1	38-39	GTCCCAGCGCTA-GCGGGCACGCGG	1.4 M b to PAK3 gene
		start codon	371-373 (Exon 4)		



Table 2i. PAK3 gene, Xq22.3

Length (nt)	SALSA MLPA probe	PAK3 exon ^a	Ligation site NM_002578.5	Partial sequence ^b (24 nt adjacent to ligation site)	Distance to next probe
Mutations in the <i>PAK3</i> gene have been reported as being the cause of non-syndromic intellectua (MRX30 or MRX47, see https://www.omim.org/entry/300558).					
		start codon	578-580 (Exon 5)		
385	02908-L03178	Exon 5	655-656	CGGGATTCTTCA-GCACTCAACCAC	40.5 kb
418	02909-L02303	Exon 10	1252-1253	CCACCCTCTGCT-GAAAATGCCAAT	52.8 kb
400	03521-L02304	Exon 17	2014-2015	ACTAATGGAACT-CCAGAGCTCCAG	4.0 kb
481	02911-L02305	Exon 18	2185-2184 reverse	TTAATTGCTTCC-TTTGCAGCGATA	113 kb to DCX gene
		stop codon	2210-2212 (Exon 18)		

Table 2j. DCX gene, Xq22.3 – q23

Length (nt)	SALSA MLPA probe	DCX exon ^a	Ligation site NM_178152.3	Partial sequence ^b (24 nt adjacent to ligation site)	Distance to next probe	
disabili https://	Mutations in the <i>DCX</i> gene are found to result in lissencephaly ('smooth brain'), characterised by intellectual disability and seizures. For more information on <i>DCX</i> -related disorders see https://www.ncbi.nlm.nih.gov/books/NBK1185/.					
P061 L	issencephaly: co	ontains more	probes for the DCX ge	ne.		
		start codon	1173-1175 (Exon 7)			
275	04124-L03481	Exon 4	824-825	GATGATGTGTTT-ATTGCCTGTGGT	67.9 kb	
215	04123-L15828	Exon 3	716-717	GTCCTCACTGAT-ATCACAGAAGCC	9.8 kb	
319	04121-L08390	Exon 2	NM_178152.3; 431 nt before exon 2; NM_000555.3; 295- 296	CAGGCTATGGAT-TCATTTACAACT	4.7 M b to AGTR2 gene	
		stop codon	78-80 (Exon 2)			

Table 2k. AGTR2 gene, Xq23

Length (nt)	SALSA MLPA probe	AGTR2 exonª	Ligation site NM_000686.5	Partial sequence ^b (24 nt adjacent to ligation site)	Distance to next probe
Mutations in the <i>AGTR2</i> gene have been reported as being the cause of non-syndromic intellectual dis (MRX88, see https://www.omim.org/entry/300852), often accompanied by seizures.					tual disability
,		start codon	169-171 (Exon 3)		
355	02925-L02319	Exon 1	31-32	TGAGAGAACGAG-TAAGCACAGAAT	2.1 kb
130	13917-L02320	Exon 3	773-774	TTTCCCACCTGA-GAAATATGCCCA	20.4 M b to ARHGEF6 gene
		stop codon	1258-1260 (Exon 3)		

Table 2I. ARHGEF6 gene, Xq26.3

Length (nt)	SALSA MLPA probe	ARHGEF6 exonª	Ligation site NM_004840.3	Partial sequence ^b (24 nt adjacent to ligation site)	Distance to next probe
	Mutations in the ARHGEF6 gene have been reported as being the cause of non-syndr disability (MRX46, https://www.omim.org/entry/300436).				c intellectual
		start codon	2379-2381 (Exon 22)		
202	02902-L04460	Exon 19	2027-2028	GATGCTCAAATC-CTTAAAGTGATC	31.9 kb
172	16857-L19651	Exon 9	1002-1003	ACAAAGTAGGAG-GTTGTCTACTGA	38.3 kb
443	16856-L19650	Exon 4	417-418	GTGGACGTTCCT-CTTCTCTTAGTG	35.5 kb
343	22017-L02293	Exon 1	120-121	CTAAAAAGACCA-TCTGTGATCCGG	11.1 M b to <i>FMR1</i> gene
		stop codon	51-53 (Exon 1)		



Table 2m. FMR1 gene, Xq27.3

Length	SALSA MLPA	FMR1	Ligation site	Partial sequence ^b	Distance to		
(nt)	probe	exonª	NM_002024.6	(24 nt adjacent to ligation site)	next probe		
Defect	Defects in the <i>FMR1</i> gene, result in fragile X syndrome, characterised by moderate to severe intellectual						

disability. Expansion of a trinucleotide repeat in exon 1 of the *FMR1* gene is the most common defect of this gene. This expansion can result in silencing of the gene due to methylation of the promoter sequence. For more information on *FMR1* disorders see https://www.ncbi.nlm.nih.gov/books/NBK1384/.

ME029 FMR1/AFF2: this methylation-specific probemix contains more probes for the *FMR1* gene and allows detection of both copy number changes, as well as the detection of promoter methylation (in full mutation male samples) of the *FMR1* and *AFF2* genes. It is not possible to directly measure the length of the trinucleotide repeat by MLPA

		start codon	262-264 (Exon 1)		
166	02927-L03721	Exon 9	1091-1092	AAAAGCTAGAAG-CTTTCTCGAATT	12.9 kb
142	02928-L03720	Exon 16	1939-1940	ACTCCCGAACAG-ATAATCGTCCAC	556 kb to AFF2 gene
		stop codon	2158-2160 (Exon 17)		

Table 2n. AFF2 gene, Xq28

Length (nt)	SALSA MLPA probe	AFF2 exon ^a	Ligation site NM_002025.4	Partial sequence ^b (24 nt adjacent to ligation site)	Distance to next probe	
FMR1, Inactiv https:// ME029 detecti	The long <i>AFF2</i> gene is located at close distance (550 kb) from <i>FMR1</i> and spans almost 500 kb. Similar to <i>FMR1</i> , expansion of a trinucleotide repeat in exon 1 of the <i>AFF2</i> gene can result in inactivation of the gene. Inactivation of the <i>AFF2</i> gene has been associated with intellectual disability (FRAXE, see https://www.omim.org/entry/309548), premature ovarian failure and obsessive-compulsive disorder. ME029 FMR1/AFF2: this methylation-specific probemix contains more probes for the <i>AFF2</i> gene and allows detection of both copy number changes, as well as the detection of promoter methylation of the <i>AFF2</i> gene. It is not possible to directly measure the length of the trinucleotide repeat by MLPA.					
		start codon	482-484 (Exon 1)			
154 «	03511-L04202	Exon 1	503-504	TCGACTTTTTCA-GAGACTGGGACT	161.1 kb	
283	00493-L00066	Exon 3	980-981	GTCATAACCCTA-GCACTGTACTGG	175.5 kb	
241	03516-L15823	Exon 5	1606-1607	CTCACTTCCATG-CATACTGCTGGA	118.4 kb	
337	02932-L02323	Exon 11	2539-2540	GAACCAAGACCT-AACATCCCTTTG	31.3 kb	
263	263 02933-L23673 Exon 20 4130-4131 CAGTGTCTCTCA-ACAACGTCTCCC					
		stop codon	4415-4417 (Exon 21)			

Table 2o. SLC6A8 gene, Xq28

Length (nt)	SALSA MLPA probe	SLC6A8 exonª	Ligation site NM_005629.4	Partial sequence ^b (24 nt adjacent to ligation site)	Distance to next probe		
by intell	Mutations in the <i>SLC6A8</i> gene are reported to cause cerebral creatine deficiency syndrome 1 characterised by intellectual disability (see https://www.omim.org/entry/300352). For more information on creatine deficiency syndromes see https://www.omim.org/entry/300352). For more information on creatine deficiency syndromes see https://www.ncbi.nlm.nih.gov/books/NBK3794/ .						
P049 SL	C6A8 - ABCD1: 0	contains mor	e probes for the SLC6A	18 gene.			
		start codon	650-652 (Exon 1)				
208 « #	01871-L15827	Exon 3	1224-1225	AGACTGTGCCAA-TGCCAGCCTGGC	3.7 kb		
188 « #	# 22782-L32127 Exon 13 2555-2554 reverse GCTGAGTTGTCA-CATGACACTCTC				705 kb to GDI1 gene		
		stop codon	2555-2557 (Exon 13)				

Table 2p. *GDI1* gene, Xq28

Length (nt)	SALSA MLPA probe	GDI1 exon ^a	Ligat NM_0	Ligation sitePartial sNM_001493.3(24 nt adjacen		Partial sequence ^b (24 nt adjacent to ligation site)				Distance to next probe	
	ons in the <i>GL</i> /www.omim.org			X-linked	intellectual	disability	(MRX41	or	MRX48,	see	

Length (nt)	SALSA MLPA probe	GDI1 exon ^a	Ligation site NM_001493.3	Partial sequence ^b (24 nt adjacent to ligation site)	Distance to next probe
		start codon	102-104 (Exon 1)		
137 «	16875-L19669	Exon 1	106-107	CCTGACCATGGA-CGAGGAATACGA	3.9 kb
378 «	16874-L23559	Exon 7	876-877	TGGATGACATCA-TCATGGAGAACG	
		stop codon	1443-1445 (Exon 11)		

^a See section



Exon numbering on page 2 for more information.

^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.

« 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.

This probe's specificity relies on a single nucleotide difference compared to a related gene or pseudogene. As a result, an apparent duplication of only this probe can be the result of a non-significant single nucleotide sequence change in the related gene or pseudogene.

~ More variable. This probe has been reported to be deleted/duplicated in healthy individuals (various reports).

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.

Complete probe sequences are available at www.mrcholland.com.

Related SALSA MLPA probemixes

- P245 / P064 Microdeletion Syndromes-1A / Microdeletion Syndromes-1B. Probes are included for different microdeletion syndromes.
- P036 / P070 Subtelomeres Mix 1 / Subtelomeres Mix 2B. Each probemix contains one probe for every subtelomere.

Several syndrome-specific probemixes useful for confirmation of findings with this X-linked ID probemix are available; see Table 2.

References

- Ropers HH et al. (2005). X-linked mental retardation. Nat Rev Genet. 6:46-57.
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Selected publications using SALSA MLPA Probemix P106 X-linked ID

- Bogliş A et al. (2020). Exon 21 deletion in the *OPHN1* gene in a family with syndromic X-linked intellectual disability. *Medicine*. 99:e21632.
- Neira VA et al. (2013). *De novo MECP2* disomy in a Mexican male carrying a supernumerary marker chromosome and no typical Lubs syndrome features. *Gene*. 524:381-385.
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P106 product history	
Version	Modification
D2	The length of one probe has been adjusted.
D1	Two probes have been replaced. Three probes have been changed in length, not in sequence detected.
C1	One ARX probe has been replaced and the 118 nt Y probe has been elongated to 121 nt.
B2	One RPS6KA3 probe has been removed and the Y-chromosome fragment on 118 nt and the control fragments (QDX2) have been replaced.
B1	Two probes for the <i>HUWE1</i> gene and one extra AGTR2 probe have been included. In addition, two ARX probes and one SLC6A8 probe have been replaced. Finally, extra control fragments at 88-96-100 and 105 nt have been included.
A1	First release.

Implemented changes in the product description

Version D2-02 - 19 November 2024 (04P)

- The term 'mental retardation' is considered outdated and was updated to 'intellectual disability' where appropriate.

Version D2-01 – 27 September 2022 (04P)

- Product description rewritten and adapted to a new template.

- Product description adapted to a new product version (version number changed, changes in Table 1 and Table 2).

- Various minor textual or layout changes.

- Small changes of probe lengths in Table 1 and 2 in order to better reflect the true lengths of the amplification products.

Version D1-01 – 21 October 2020 (02P)

- Product renamed from "MRX" to "X-linked ID".

- "Intellectual disability" has replaced the term "mental retardation" throughout this document.

- Product description rewritten and adapted to a new template.

- Product description adapted to a new product version (version number changed, changes in Table 1 and Table 2).

- Various minor textual or layout changes.

- Ligation sites of the probes targeting all genes except for *PQBP1* updated according to new version of the NM_ reference sequence.

- Warning added to Table 1 and 2 for probe specificity relying on a single nucleotide difference between target gene and related gene or pseudogene.

- For uniformity, the chromosomal locations and bands in this document are now all based on hg18 (NCBI36).

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