

Product Description

SALSA® MLPA® Probemix P520-A2 MPN mix 2

To be used with the MLPA General Protocol.

Version A2

As compared to version A1, several probes have been changed in length, but not in the sequence detected. For complete product history see page 11.

The P520-A2 MPN mix 2 assay allows mutations detection with high sensitivity. Only 1% allele burden is required for detection of the eight different mutations frequently found in myeloproliferative neoplasms (MPNs).

With each P520 probemix order, an artificial DNA sample (SD057) with an estimated 1% allele burden is supplied for data binning in the fragment analysis procedure, and for serving as an artificial positive control for all the mutation-specific probes. This SD057 should be used in each P520 experiment for correct fragment binning and mutation calling.

Please note that this probemix should be used only as a tool for determining mutation presence (or absence) and NOT used as a quantitative tool for measuring exact mutation burden. In addition, positive controls should be included both in the validation phase and in experiments.

Catalogue numbers:

- P520-025R: SALSA MLPA Probemix P520 MPN mix 2, 25 reactions.
- P520-050R: SALSA MLPA Probemix P520 MPN mix 2, 50 reactions.
- P520-100R: SALSA MLPA Probemix P520 MPN mix 2, 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.

Note 1: The use of this product requires two modifications to the MLPA General Protocol. First, the PCR reaction should be prolonged from 35 to 40 cycles. Second, we strongly recommend to use ≥100 ng of DNA sample per MLPA reaction (maximum 400 ng of DNA) for most optimal and robust mutation detection with the P520 probemix. P520 probemix users are strongly advised not to make any additional modifications to the MLPA protocol, as this could potentially lead to false positive or false negative results for the mutation calling.

Note 2: Probemix P420 MPN mix 1 can be used for test samples with a >10% allele burden, which will generate a saturated mutation-specific signal when using P520.

This MLPA probemix cannot detect any mutations that are outside the target sequences of the MLPA probes. Neither can it be used for copy number detection of the *JAK2*, *MPL*, *CALR* and *KIT* target genes. Even when MLPA analysis did not detect any aberrations, the possibility remains that changes in the corresponding gene(s) do exist but remain undetected. This probemix covers only the most frequent mutations of clinical and diagnostic relevance in MPNs. Rare or infrequent mutations are not covered. **If mutation calling with**



MLPA is negative for a test sample, we recommend to perform sequencing of the *JAK2*, *MPL*, *CALR* and *KIT* genes to detect the rare or unknown mutations possibly present in a test sample.

Please note that the probe signals of the reference probes in this P520 probemix show much greater variation as compared to standard MLPA products. This is due to the nature of the high sensitivity of this assay.

General information

The SALSA MLPA Probemix P520 MPN mix 2 is a **research use only (RUO)** assay for detection of eight different mutations frequently found in MPNs in *JAK2*, *MPL*, *CALR* and *KIT* genes.

MPNs are clonal hematopoietic stem cell malignancies, characterized by excessive production of blood cells. MPNs are subdivided in polycytemia vera (PV), essential thrombocytemia (ET), primary myelofibrosis (PMF) and less common conditions like chronic neutrophilic leukemia (CNL), chronic eosinophilic leukemia (CEL), hypereosinophilic syndrome (HES) and mastocytosis.

Discovery of a frequent *JAK2* mutation (9p24.1), common to classic MPNs (PV, ET and PMF), has linked these diseases on a molecular level. The current WHO diagnostic criteria for classic MPNs include presence of *JAK2*, *CALR* or *MPL* mutations. The *JAK2* V617F point mutation is detected in ~98% of PV patients, and in ~60% of patients with ET and PMF, whereas other *JAK2* exon 12 mutations are commonly found in V617F negative PV patients. P520-A2 MPN mix 2 contains three mutation-specific *JAK2* probes: one probe for V617F and two probes for the most common exon 12 mutations N542_E543del and E543_D544del.

Mutations in the *MPL* gene (1p34.2) are found in 4-11% of *JAK2* V617F negative ET and PMF patients. This problemix contains two mutation-specific probes for *MPL*, W515K and W515L, that are diagnostically relevant in PV, ET and PMF according to the WHO classification.

The discovery of novel *CALR* gene (19p13.13) mutations in ET and PMF provides additional diagnostic tools for MPNs. Patients with ET and PMF but negative for *JAK2* and *MPL* mutations, have been reported to harbour somatic insertions and deletions in exon 9 of the *CALR* gene. A 52-bp deletion (type 1) and a 5-bp insertion (type 2) are the most common mutations found in the *CALR* gene (53% and 32%, respectively). These mutations result in a frameshift to an alternative reading frame (Klampfl et al. 2013, Nangalia et al. 2013). *CALR* mutation-specific probes for the 52-bp deletion (L367fs*46, type 1) and 5-bp insertion (K385fs*47, type 2) are included in this probemix.

In addition, a probe specific for the D816V mutation in the *KIT* gene (4q12) is present. This is the most common *KIT* mutation and is present in >90% of patients with systemic mastocytosis (SM). Consequently, the presence of this mutation is considered a diagnostic criterion of SM according to the WHO classification.

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 JAK2, MPL, CALR and KIT exon numbering used in this P520-A2 MPN mix 2 product description is the exon numbering from the LRG_612 (JAK2), LRG_510 (MPL), LRG_828 (CALR) and LRG_307 (KIT). 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 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 P520-A2 MPN mix 2 contains 25 MLPA probes with amplification products between 115 and 338 nucleotides (nt). This probemix contains eight probes specific for the *JAK2*, *MPL*, *CALR*





and *KIT* mutations which will only generate a signal when the mutation is present (see details of these mutations detected below).

JAK2 p.V617F = c.1849G>T

JAK2 p.N542_E543del = c.1624_1629delAATGAA

JAK2 p.E543_D544del = c.1627_1632delGAAGAT

ARD: p.W515K = c.1542_1544TC>AA

 $MPL p.W515K = c.1543_1544TG>AA$

MPL p.W515L = c.1544G>T

CALR p.L367fs*46 = c.1092_1143del52 **CALR** p.K385fs*47 = c.1154_1155insTTGTC

KIT p.D816V = c.2447A>T

This probemix is modified to allow higher detection sensitivity for mutations. Only >1% allele burden is needed for reliable detection of the above mentioned mutations.

Finally, 17 reference probes are included in P520-A2 probemix, detecting different autosomal chromosomal locations which are relatively stable in MPNs. Complete probe sequences and the identity of the genes detected by the reference probes are available online (www.mrcholland.com).

In mutation-negative samples, only the 17 reference probes are expected to generate a signal. The remaining eight probes will only generate a signal when the corresponding mutation is present in the test sample. Please note that, however, mutation-specific probes can generate a low background signal in normal samples.

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, and specifically for this probemix in Figure 1 of this product description.

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 the MLPA technique (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).

Modifications to the MLPA General Protocol (One-Tube)

Please note that this P520 probemix should be used together with a modified MLPA General Protocol with 40 PCR cycles. In addition, we recommend to use 100 ng of DNA as a minimum amount of sample DNA per MLPA reaction. P520 probemix users are strongly advised not to make any additional modifications to the MLPA protocol, as this could potentially lead to false positive or false negative results for the mutation calling. The standard One-Tube MLPA protocol should be followed, but using the following thermocycler program for the MLPA reaction:





DNA dens	DNA denaturation				
DIVA della			F		
	1.	98°C	5 minutes		
	2.	25°C	pause		
Hybridisa	tion rea	action			
	3.	95°C	1 minute		
	4.	60°C	16-20 hours		
Ligation r	reaction	1			
	5.	54°C	Pause		
	6.	54°C	15 minutes		
	7.	98°C	5 minutes		
	8.	20°C	pause		
PCR reac	tion				
	9.	40 cycles:	95°C	30 seconds	
			60°C	30 seconds	
			72°C	60 seconds	
	10.	72°C	20 minutes		
	11.	15°C	pause		

MLPA technique validation

Internal validation of the MLPA technique using 16 DNA samples from healthy individuals 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 \leq 0.20 for all reference 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 (≥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 healthy individuals without a history of myeloproliferative neoplasms. More information regarding the selection and use of reference samples can be found in the MLPA General Protocol (www.mrcholland.com).

Positive and negative control DNA samples

MRC Holland cannot provide positive DNA samples. Inclusion of a positive sample in each experiment is recommended. The quality of cell lines can change; therefore samples should be validated before use. Please find below some sources for commercial positive reference samples, tested by MRC Holland:

- Reference standard sample for the *JAK2* p.V617F point mutation (Catalog IDs: HD649 for mutant and HD652 for WT) available at www.horizondiscovery.com (Horizon Discovery).
- JAK2 p.V617F WHO reference panel (code 16/120) (nominal values 100% [100%], 90% [89.5%], 30% [29.6%], 10% [10.8%], 1% [1.0%], 0.1% [0.03%] and 0% [0%] VAF in square brackets is the proposed consensus value by WHO committee) www.nibsc.org (The National Institute for Biological Standards and Control NIBSC).
- Quan-Plex™ NGS Reference Standard Genomic DNA (Catalog ID: ARF-1001G-1 for mutant) including KIT p.D816V and Onco-Ref™ Genomic DNA Reference Standard HCT116 WT (Catalog ID: ASO-6052-1) www.accuref.com (AccuRef).

We highly recommend inclusion of **negative control samples** from healthy individuals in the experiments to evaluate the background signal for the mutation-specific probes in the MLPA peak pattern for your specific types of samples and DNA extraction method used.





Additionally, we highly recommend to also include mutation positive samples (cell line or patient derived) extracted with the same DNA extraction method as the test samples in MLPA experiments to optimise mutation calling for the test samples.

SALSA Binning DNA SD057

The SD057 Binning DNA provided with this probemix can aid the binning and serve as a mutation calling threshold sample for the eight mutation-specific probes: CALR L367fs*46 (S0999-L26702; 124 nt), CALR K385fs*47 (S1001-L26517; 130 nt), JAK2 N542_E543del (16924-L21237; 167 nt), JAK2 E543_D544del (16924-L21238; 172 nt), MPL W515K (S1048-SP0405-L29870; 181 nt), MPL W515L (S1048-SP0405-L29871; 186 nt), KIT D816V (17722-SP0542-L23707; 200 nt) and JAK2 V617F (13190-L21572; 240 nt). SD057 Binning DNA is a mixture of genomic DNA from healthy individuals and plasmid DNA that contains the target sequence detected by the above mentioned probe. NOTE that this SD057 sample DNA contains an estimated 1% burden of the above mentioned mutation-specific sequences. Inclusion of one reaction with 5 μl SD057 (20 ng/μl) Binning DNA with total of 100 ng of DNA in each MLPA experiment is essential as it can be used to aid in data binning of the peak pattern using Coffalyser.Net software and as an artificial positive control for the specific point mutations. For reliable threshold determination and subsequently mutation calling, it is highly recommended to include three reactions of SD057 in each MLPA experiment (please see Interpretation of results for mutation calls section for more information). Furthermore, Binning DNA should be included in the experiment whenever changes have been applied to the set-up of the capillary electrophoresis device (e.g. when capillaries have been renewed). It is strongly advised that all samples tested are extracted with the same method and derived from the same source of tissue. For further details, please consult the SD057 Binning DNA product description, available online: www.mrcholland.com. This product is for research use only (RUO).

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.

In contrast to conventional MLPA probemixes, the Q-fragments in P520 will provide a warning in Coffalyser.Net already when less than 100 ng of sample DNA is used per reaction. See Figure 1 below for the effect of DNA quantity on the Q-fragments. The more sample DNA is used per reaction, the lower the Q-fragments. Q-fragments are high when DNA amount is too low or ligation has failed. When all four Q-fragment have signals >33% of the 92 nt control fragment this indicates that DNA quantity was insufficient for the MLPA reaction (<100 ng) and Coffalyser.Net software will indicate a warning for that particular sample. Additional information on how to interpret observations on the D/X/Y-control fragments can be found in the MLPA protocol.

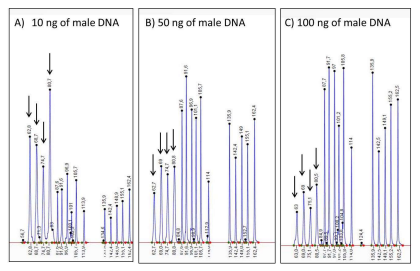


Figure 1. Q-fragments in the P520-A2 **probemix.** P520-A2 probemix analysed with A) 10 ng of male DNA; median of Qfragments is higher than the 92 nt control fragment and in Coffalyser.Net analysis a red warning is indicated; B) 50 ng of male DNA; median of Q-fragments is higher than 33% of the 92 nt control fragment and in Coffalyser.Net an orange warning is indicated; C) 100 ng of male DNA, Qfragments are <33% of the 92 nt probe and no warning is indicated in Coffalyser.Net). The Q-fragments are indicated with arrows. Note that the MLPA peak pattern of the P520-A2 probemix is shown here only partially, until 162 nt.





Interpretation of results for mutation calls

The standard deviation of each individual reference probe over all the reference samples should be ≤ 0.20 . Please note that Coffalyser.Net uses stricter cut-off for standard deviation, which is not applicable for this probemix with the condition that the experiment includes ≥ 3 reference samples extracted from the same tissue/sample type with same pre-analytical conditions and DNA extraction method. When these criteria are fulfilled, the assay can yield reliable results for point mutation detection and calling.

Comparison of the relative peak height of a mutation-specific probe (vs. reference probes) on a positive sample with the relative peak height obtained with that probe on the SD057 Binning DNA serves as a threshold for the presence for the corresponding mutation. SD057 contains an estimated 1% allele burden for each aforementioned mutation. Given that **three reactions with SD057 are included in the experiment and defined as 'Positive reference' as 'sample type'** in *fragment analysis* tab, the intra ratio percentages in the Coffalyser 'comparative analysis experiment explorer' in *ratio overview* (for all samples) or *sample report* (per sample) tabs can be directly compared between SD057 for reliable mutation calling.

In case the signal of the mutation-specific probe in a sample of interest is higher than the average signal of the corresponding probe in the three SD057 reactions, serving as threshold samples, this is considered to be a true positive call for the respective mutation. When the mutation-specific signal is lower than the corresponding mutation-specific probe average signal in the three SD057 reactions, this is regarded as a negative call (see examples in Figure 2).

When probe signals at 167 nt and 172 nt are both present in the MLPA peak pattern, and higher than in the SD057 threshold sample, it is indicative for the *JAK2* E543_D544del mutation. When only the probe signal at 167 nt is present, and higher than in the SD057 threshold sample, it is indicative for the *JAK2* N542_E543del mutation.

In addition to Coffalyser.Net analysis, visual examination of the P520 MLPA peak patterns is always necessary for mutation calling (in addition to Figure 2 for negative mutation calling, see also Figure 4 for saturated mutation-specific MLPA peaks) to confirm that the signals seen in the sample of interest are higher than in the SD057 threshold sample(s).



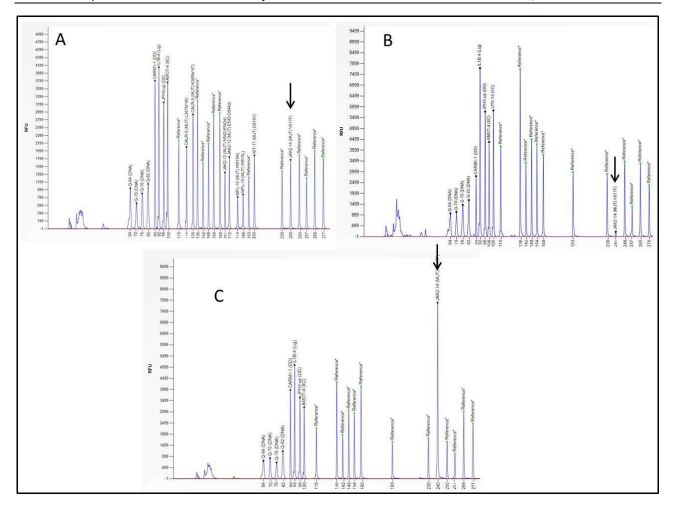


Figure 2. Mutation calling with the P520-A2 probemix: examples of negative and positive mutation calls for the *JAK2* **V617F point mutation.** A) P520-A2 probemix analysed with SD057 Binning DNA that contains an estimated 1% burden of the *JAK2* V617F mutation sequence. The peak of the *JAK2* V617F specific probe is indicated with an arrow. B) An example of a sample from a healthy individual that has a very small background peak at the length of the *JAK2* V617F specific probe (at 240 nt, indicated with an arrow). As the height of this probe is lower than the height of the *JAK2* V617F specific probe in SD057, this is considered a negative call for the *JAK2* V617F point mutation. C) An example of a test sample with 5% allelic burden for the *JAK2* V617F mutation (peak at 240 nt, indicated with an arrow). As the height of this signal is higher in comparison with the *JAK2* V617F signal in the SD057 Binning DNA (A) this results can be considered as a positive call for the *JAK2* V617F mutation. Please note that the MLPA peak pattern for the P520-A2 probemix is shown here only partially, until 276 nt for better visualisation.

Saturated MLPA peak(s) with >5% allele burden samples

When a test sample has >5% allele burden, the corresponding MLPA probe peak may become saturated in the MLPA peak pattern (see Figure 3 for further information). The saturated peak height is usually much higher than the upper threshold for relative fluorescence detection limit of capillary electrophoresis instrument.

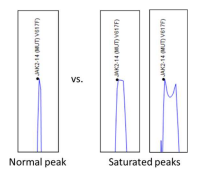


Figure 3. Normal vs. saturated MLPA peaks. Saturated peaks can be detected in the MLPA peak pattern by zooming in: a blunt end of an MLPA peak or alternatively a double peak at the very tip of the MLPA peak is then detected.



High allele burden (>5%) commonly results in an extra "shadow peak" ~15 nt before the real mutation-specific peak. This "shadow peak" can occur to any of the mutation-specific probes and it has the exact same nucleotide sequence as the real MLPA peak, however, due to very high concentration of target sequence, is forming additional secondary structures that result in a faster running speed in the capillary electrophoresis and in formation of the "shadow peak" (see Figure 4 for further information).

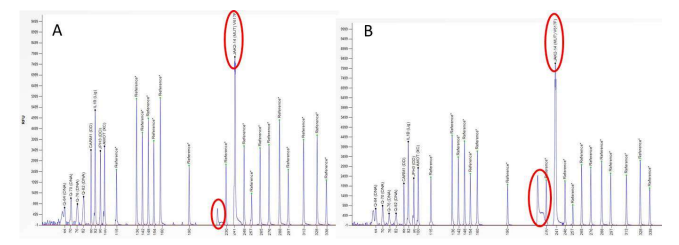


Figure 4. Saturated mutation-specific peaks in the MLPA peak pattern. A) As an example *JAK2* V617F positive test sample with 10% allele burden was analysed with the P520-A2 probemix and run at an ABI sequencer. The *JAK2* V617F specific peak at 240 nt is saturated and there is a "shadow peak" visible at 222 nt. B) The same *JAK2* V617F positive cell line sample with 50% allele burden and run at the ABI sequencer showing an increased shadow peak at 222 nt.

When a "shadow peak" (the unspecific peak ~15 nt from the saturated mutation-specific peak) is detected in the MLPA peak pattern, a rerun on the capillary electrophoresis device using a dilution (e.g. 20x) of the PCR product with pure water can be used to lower this "shadow peak". Please note that with this dilution the other probe signals in the MLPA peak pattern are diminished as well.

Please note that the final ratios and mutation calling are affected both by percentage of tumour cells and by possible subclonality.

- False positive results: Literature suggests that some individuals in the general population can have *JAK2* V617F mutation-positive clones at low levels (See e.g. Sidon et al. 2006, Nielsen et al. 2013). Please note also that abnormalities detected by a single probe 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. 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>Copy number changes detected by reference probes</u> 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. The risk of 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 softwares might 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

 This SALSA MLPA probemix P520 is designed to detect the presence of the aforementioned mutations in a DNA sample. Note that a mutation or polymorphism in the sequence detected by a probe can also cause a reduction in relative peak height, even when not located exactly on the ligation site, and this can lead to





false positive or negative results! In addition, some probe signals can be more sensitive to sample purity and small changes in experimental conditions. Unlike majority of MLPA probemixes this P520 does not provide copy number information of the target probes.

- MLPA cannot detect mutations that are not included in this assay. 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.
- 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 test sample, especially in tumour samples with more chaotic karyotypes.

Confirmation of results

Confirmation of the presence of a specific point mutation can be done by e.g. Sanger sequencing, next generation sequencing, qPCR, or digital droplet PCR. In particular, it is recommended to confirm positive mutation calls with low mutation burden with the above mentioned methods.

We recommended to use P420 MPN mix 1 for test samples when there is a saturated mutation-specific signal in P520 in case of >10 % allele burden present in the test sample and in order to confirm the result. The P420 assay is a conventional MLPA assay allowing detection of mutations on samples with >10-20% allele burden (up to 100% allele burden). For samples with estimated 1-10% allele burden we recommend to use P520 as qualitative tool for mutation detection.

COSMIC mutation database

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

Please report false positive or false negative results due to SNVs and unusual results to MRC Holland: info@mrcholland.com.





Table 1. SALSA MLPA Probemix P520-A2 MPN mix 2

Longeth (nt)	CALCA MI DA mucho	Chromosomal position (hg18) ^a		
Length (nt)	SALSA MLPA probe	Reference	Mutation	
64-105	Control fragments – see table in probemix content section for more information			
115	Reference probe S0973-L26704	4p13		
124 §	CALR probe S0999-L26702		p.L367fs*46 =c.1092_1143del52	
130 §	CALR probe S1001-L26517		p.K385fs*47 =c.1154_1155insTTGTC	
136	Reference probe 16316-L25926	3q21		
142	Reference probe 07387-L26769	12q13		
148	Reference probe 10663-L11245	6p12		
154	Reference probe 13781-L15275	11p14		
160	Reference probe 17621-L21665	10q22		
167§£	JAK2 probe 16924-L21237		p.N542_E543del=c.1624_1629delAATGAA	
172 § £	JAK2 probe 16924-L21238		p.E543_D544del=c.1627_1632delGAAGAT	
181 § Ж¥	MPL probe S1048-SP0405-L29870		p.W515K =c.1543_1544TG>AA	
186 § Ж¥	MPL probe S1048-SP0405-L29871		p.W515L =c.1544G>T	
193¥	Reference probe 11556-L26606	5q31		
200 § Ж	KIT probe 17722-SP0542-L23707		p.D816V =c.2447A>T	
230	Reference probe 17130-L26574	11p11		
240 § ∑¥	JAK2 probe 13190-L21572		p.V617F =c.1849G>T	
250 ¥	Reference probe 05386-L29196	12p11		
257¥	Reference probe 13572-L29195	1q23		
265	Reference probe 12434-L26073	14q24		
277	Reference probe 16270-L26771	20q11		
287	Reference probe 05713-L20268	2p11		
297	Reference probe 04570-L20036	16q13		
313	Reference probe 04833-L20693	5p13		
328	Reference probe 13397-L26608	6q12		
338	Reference probe 12785-L15496	2q13		

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

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.

[¥] Changed in version A2 (from lot A2-0217 onwards). Small change in length, no change in sequence detected.

[§] Mutation-specific probe. This probe will generate a signal when the mutation is present.

Ж This probe consists of three parts and has two ligation sites.

 $[\]pounds$ When probe signals at 167 nt and 172 nt are both present in the MLPA peak pattern, and the signals are higher than in the SD057 threshold sample, it is indicative for the *JAK2* E543_D544del mutation. When only the probe signal at 167 nt is present, and the signal is higher than in the SD057 threshold sample, it is indicative for the *JAK2* N542_E543del mutation.

 $[\]Sigma$ This probe can have a very low unspecific background signal detected also in healthy control samples. This background signal has never exceeded the signal detected in *JAK2* V617F samples with an 1% allele burden in the internal quality tests at MRC Holland. See Figure 2 for further information on this.





Table 2. P520-A2 probes arranged according to chromosomal location

Length (nt)	SALSA MLPA probe	Gene / exon ^a / Mutation	Ligation site	Partial sequence ^b (24 nt adjacent to ligation site)	
	MPL gene, at 1p34.2, indicated ligation sites are in NM_005373.3.				
detected in	Mutations in the thrombopoietin receptor (<i>MPL</i>) gene, most commonly W515L and W515K substitutions in exon 10, are detected in patients with ET and PMF, which can support diagnosis when <i>JAK2</i> is not mutated. W515L and W515K are reported to be the predominant <i>MPL</i> mutations in MPNs (Ma W et al. 2011).				
186§Ж	S1048-SP0405- L29871	<i>MPL</i> , exon 10, p.W515L=c.1544G>T	1575-1576 and 25 nt after exon 10	GCTGCTGATGT T -46nt spanning oligo-TGGCGGTGGACC	
181 § Ж	S1048-SP0405- L29870	MPL, exon 10, p.W515K=c.1543_1544TG>AA	1575-1576 and 25 nt after exon 10	GCTGCTGAGG AA -46nt spanning oligo-TGGCGGTGGACC	
VIT gane at 4g12 indicated ligation sites are in NM 000222.2					

KIT gene, at 4q12, indicated ligation sites are in NM_000222.3.

Somatic activating *KIT* D816V mutation is present in more than 97% of SM patient samples (Erben P et al. 2014). This D816V mutation results in ligand-independent activation of c-KIT tyrosine kinase, and detection of this point mutation aids in prediction of response to tyrosine kinase inhibitor (TKI) therapy (imatinib).

200 S W	17722-SP0542-	<i>KIT</i> , exon 17 ,	2505-2504 and	CATTCTTGATGA-38nt spanning
200§Ж	L23707	p.D816V=c.2447A>T	2467-2466 reverse	oligo-CCATGAGTAAGG

JAK2 gene, at 9q24.1, indicated ligation sites are in NM_004972.4.

Discovery of a JAK2 mutation common to MF, PV and ET has linked these diseases on a molecular level, and diagnostic criteria for MPN include detection of a clonal marker e.g. JAK2 V617F mutation in exon 14 or exon 12 mutations. JAK2 is mutated in 97% of all patients with PV, 55% of patients with ET and 65% of patients with PMF (Tefferi A, 2010). The most common JAK2 mutations in exon 12 are the following 6-bp deletions: N542_E543del and E543_D544del.

167§£	16924-L21237	JAK2, exon 12, p.N542_E543del =c.1624_1629delAATGAA	2099-2100	AAAATCAGAGAT-TTGATATTTGTA
172 § £	16924-L21238	JAK2, exon 12, p.E543_D544del =c.1627_1632delGAAGAT	2099-2100	AAAATCAGAAAT-TTGATATTTGTA
240 § ∑	13190-L21572	JAK2, exon 14, p.V617F=c.1849G>T	2316-2315 reverse	GTCTCCACAGA A -ACATACTCCATA

CALR gene, at 19p13.13, indicated ligation sites are in NM_004343.4.

Majority of patients with ET or PM that are negative for *JAK2* and *MPL* carry a somatic mutation in *CALR* gene. A 52-bp deletion (type 1=L367fs*46) and 5-bp insertion (type 2=K385fs*47) are the most common mutations found in *CALR* (53% and 32%, respectively), resulting in a frameshift to an alternative reading frame (Klampfl T et al. 2013, Nangalia J et al. 2013).

130 §	S1001-L26517	CALR, exon 9, p.K385fs*47 =c.1154_1155insTTGTC	1225-1226	CAGAGGACAA TT-GTC GGAGGATGA
124§	S0999-L26702	CALR, exon 9, p.L367fs*46 =c.1092_1143del52	1163-1214 reverse	сттетсстстес-тестсетсстет

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

§ Mutation-specific probe. This probe will generate a signal when the mutation is present.

Ж This probe consists of three parts and has two ligation sites.

£ When probe signals at 167 nt and 172 nt both are present in the MLPA peak pattern, and signals are higher than in the SD057 threshold sample, it is indicative for the JAK2 E543_D544del mutation. When only the probe signal at 167 nt is present, and the signal is higher than in the SD057 threshold sample, it is indicative for the JAK2 N542_E543del mutation. ∑ This probe can have a very low unspecific background signal detected also in healthy control samples. This background signal has never exceeded the signal detected in JAK2 V617F samples with an 1% allele burden in the internal quality tests at MRC-Holland. See Figure 2 for further information on this.

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.

^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.



Related SALSA MLPA probemixes

P420 MPN mix 1	This probemix is a conventional MLPA probemix allowing detection of the same mutations
F420 WIFIN IIIIX I	as included in P520 on samples with >10-20% allele burden (up to 100% allele burden).
	This probemix contains probes for chromosome 3, 5q (EGR1, MIR145, SPARC, MIR146A),
P414 MDS	7q (EZH2), 8q (MYC), 11q (KMT2A), 12p (ETV6), chromosome 17 (TP53, NF1, SUZ12),
	chromosome 19, 20q (ASXL1), Y-chromosome, and for JAK2 V617F mutation.
P377	This probemix contains probes for 2p (MYCN, ALK), 5q (MIR145, EBF1, MIR146A), 6q, 7p12
Hematologic	(IKZF1), 7q, 8q24 (MYC), 9p (MTAP, CDKN2A, CDKN2B, PAX5), 10q23 (PTEN), 11q22 (ATM),
malignancies	12p13 (ETV6, CCND2, MDM2), 12q, 13q14 (RB1, MIR15A, DLEU2, DLEU1), 17p13 (TP53),
manynancies	17q, Chr 18, Chr 19, 21q22 (<i>RUNX1</i>) and for <i>JAK2</i> V617F mutation.

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P520 product history		
Version	Version Modification	
A2	Several probes have been changed in length, but not in the sequence detected.	
A1	First release.	

Implemented changes in the product description

Version A2-05 - 3 May 2023

- Added a sentence on page 6 that Coffalyser.net uses stricter cut-off for standard deviation, which is not applicable for this probemix when the experiment includes ≥3 reference samples.

Version A2-04 - 23 March 2023

- Correction for the link in included on page 4.
- Added in SD057 information that for threshold purposes for reliable mutation detection and calling, it is highly recommended to include three reactions of SD057 in each MLPA experiment.





- Various clarifications in the Interpretation of results for mutation calls -section on pages 6, including Coffalyser analysis instructions for mutation calling.
- Minor clarifications and adjustments throughout the document.

Version A2-03 - 13 July 2022 (04P)

- Text about SD057 on page 5 is corrected: SD057 should be used in each P520 experiment, not only for the initial experiments.

Version A2-02 - 16 December 2021 (04P)

- Modified MLPA General Protocol (One-Tube) corrected on page 4: Hybridisation reaction step 60°C is for 16-20 hours.

Version A2-01 - 11 March 2021 (04P)

- Product description adapted to a new template.
- Various minor textual or layout changes.
- Ligation sites of the probes targeting the *JAK2*, *MPL*, *CALR* and *KIT* genes are updated according to new version of the NM_ reference sequence.
- Small changes of probe lengths in Table 1 and 2 in order to better reflect the true lengths of the amplification products.

Version 09 - 26 August 2019 (T08)

- Information regarding *JAK2* p.V617F WHO reference panel has been elaborated by including the proposed consensus values and a sample for 10% VAF on page 6.

Version 08 - 09 July 2019 (T08)

- Information regarding the estimated mutation burden of SD057 has been changed back to 1% upon the release of the improved SD057-S02.
- Sources for commercial positive reference samples updated on page 6.
- New picture added to page 10 on the CE pattern of SD057-S02 using P520-A2-0217.

Version 07 - 18 October 2018 (T08)

- Information regarding the estimated mutation burden of SD057 has been changed from 1% to 0.5-1%.
- Product description modified so that P520 probemix with SD057 should be used only as a qualitative tool (mutation present or absent) and not as a quantitative tool for measuring exact mutation burden.
- Remark added that internal or external reference standards should be included both in the validation phase and in experiments.
- Information added on page 3 about possibility of detecting mutations at low level in general population.
- Recommendation about confirmation of positive mutation calls with low mutation burden and possible methods used for this purpose added on page 5.
- Information regarding positive reference samples added and adjusted on page 6.
- Various minor textual changes throughout the document.

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