

sebia

CAPI 3 HEMOGLOBIN(E)

Ref. 2507

PHORESIS VS ≥ 9.15

IVD

CE

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INTENDED USE

The CAPI 3 HEMOGLOBIN(E) kit is designed for the separation of the normal hemoglobins (A, A2 and F) in human blood samples, and for the detection of the major hemoglobin variants (S, C, E and D), by capillary electrophoresis in alkaline buffer (pH 9.4) with the SEBIA CAPILLARYS 3 instrument.

The CAPILLARYS 3 instrument is an automated analyzer which performs a complete hemoglobin profile for the quantitative analysis of the normal hemoglobin fractions A, A2 and F and for the detection of major hemoglobin variants S, C, E and D. The assay is performed on the hemolysate of whole blood samples collected in tubes containing KEDTA as anticoagulant.

For *In Vitro* Diagnostic Use.

NOTE : In this instruction sheet, the name "CAPILLARYS 3" is used for the SEBIA CAPILLARYS 3 OCTA and CAPILLARYS 3 TERA automated instruments.

PRINCIPLE OF THE TEST¹⁻²⁰

Hemoglobin is a complex molecule composed of two pairs of polypeptide chains. Each chain is linked to the heme, a tetrapyrrolic nucleus (porphyrin) which chelates an iron atom. The heme part is common to all hemoglobins and their variants. The type of hemoglobin is determined by the protein part called globin. Polypeptide chains α , β , δ and γ constitute the normal human hemoglobins :

- hemoglobin A = $\alpha_2 \beta_2$
- hemoglobin A₂ = $\alpha_2 \delta_2$
- fetal hemoglobin F = $\alpha_2 \gamma_2$

The α -chain is common to these three hemoglobins.

The hemoglobin spatial structure and other molecular properties (like that of all proteins) depend on the nature and the sequence of the amino acids constituting the chains. Substitution of amino acids by mutation is responsible for formation of hemoglobin variants which have different surface charge and consequently different electrophoretic mobilities, which also depend on the pH and ionic strength of the buffer.

The resulting qualitative (or structural) abnormalities are called hemoglobinopathies (9,10,13). Decreased synthesis of one of the hemoglobin chains leads to quantitative (or regulation) abnormalities, called thalassemias.

Hemoglobin electrophoresis is a well established technique routinely used in clinical laboratories for screening samples for hemoglobin abnormalities (1,2,3,4,12). Besides the electrophoresis techniques performed on different media, including agarose gel and chromatography, capillary electrophoresis has been developed to provide complete automation with fast separation and good resolution. It is defined as an electrokinetic separation technique carried out in a tube of internal diameter lower than 100 μm filled with a buffer composed of electrolytes. In many aspects, the methodology can be considered as an intermediary type of technique between classical zone electrophoresis and liquid chromatography (8,11).

The CAPILLARYS 3 instrument uses the principle of capillary electrophoresis in free solution which is the most common form of capillary electrophoresis. With this technique, charged molecules are separated by their electrophoretic mobility in an alkaline buffer with a specific pH. Separation also occurs according to the electrolyte pH and electroosmotic flow (5).

The CAPILLARYS 3 instrument has silica capillaries functioning in parallel allowing 8 simultaneous analyses (CAPILLARYS 3 OCTA) or 12 simultaneous analyses (CAPILLARYS 3 TERA) for hemoglobin quantification in a whole blood sample. A sample dilution with hemolysing solution is prepared and injected by aspiration at the anodic end of the capillary. A high voltage protein separation is then performed and direct detection of the hemoglobins is made at the cathodic end of the capillary at 415 nm, which is the absorbance wave length specific to hemoglobins. Before each run, the capillaries are washed with a wash solution and prepared for the next analysis with buffer.

Direct detection provides accurate relative quantification of individual hemoglobin fraction, with particular interest, such as A2 hemoglobin for β thalassemia diagnostic and the resulting electrophoregrams are also evaluated visually for pattern abnormalities. In addition, the high resolution of this procedure should allow the identification of hemoglobin variants, in particular, to differentiate hemoglobins S from D, and E from C.

The hemoglobin A2 quantification can also be performed when hemoglobin E is present.

By using alkaline pH buffer, normal and abnormal (or variant) hemoglobins are detected in the following order, from cathode to anode: $\delta\text{A}2$ (A2 variant), C, A2/O-Arab, E, S, D, G-Philadelphia, F, A, Hope, Bart's, J, N-Baltimore and H.

The carbonic anhydrase is not visualized on the hemoglobin electrophoretic patterns by capillary electrophoresis, this permits to identify hemoglobin A2 variants in this migration zone.

REAGENTS AND MATERIALS SUPPLIED IN THE CAPI 3 HEMOGLOBIN(E) KIT

WARNING : See the safety data sheets.

ITEMS	PN 2507
Buffer (ready to use)	2 vials, 700 mL each
Hemolysing solution (ready to use)	1 vial, 700 mL
Filters	4 filters

During transportation, the kit can be kept without refrigeration (15 to 30 °C) for 15 days without any adverse effects on performance.

FOR OPTIMAL MANAGEMENT OF TRACEABILITY : All reagents from the same kit must be used together.

TO OBTAIN THE EXPECTED PERFORMANCES : The package insert instructions must be observed.

WARNING : Do not use marketed deionized water, such as water for ironing for example (risk of important capillaries damage). Use only water with ultrapure quality, such as injection grade water.

1. BUFFER

Preparation

The buffer is ready to use. It contains : buffer solution pH 9.4 ± 0.5 ; additives, nonhazardous at concentrations used, necessary for optimum performance.

Use

Buffer for analysis of hemoglobins with capillary electrophoresis.

Storage, stability and signs of deterioration

Store the buffer refrigerated (2 to 8 °C). It is stable until the expiration date indicated on the kit package or buffer vial labels. Avoid storage at room temperature (15 to 30 °C) for a long time or close to a window or to a heat source.

DO NOT FREEZE.

IMPORTANT : When stored at 2 - 8 °C and prior to use, it is necessary for the buffer to reach room temperature (15 to 30 °C) ; when it is full, let the buffer vial at room temperature for at least 3 hours prior to use. If this precaution is not respected, the performances of the procedure may be affected.

WARNING : *Do not pre-heat the buffer in hot water.*

Once the buffer vial has been opened and positioned on the CAPILLARYS 3 instrument, it is stable for a maximum of **1 month** (accumulated) at room temperature (15 to 30 °C).

After each use, the buffer must imperatively be stored refrigerated (between 2 and 8 °C) without any delay, it is then stable until the expiration date indicated on the buffer vial label.

IMPORTANT : The accumulated time of the buffer stored at room temperature (15 to 30 °C) must not exceed **1 month**. This time of 1 month storage takes account of the time for the buffer to come to room temperature.

Discard buffer if it changes its appearance, e.g., becomes cloudy due to microbial contamination.

2. HEMOLYSING SOLUTION

Preparation

Hemolysing solution is ready to use. It contains buffer solution pH 8.5 ± 0.5 ; additives, nonhazardous at concentrations used, necessary for optimum performance.

Use

To dilute and hemolyze red blood cells from whole blood.

Storage, stability and signs of deterioration

Store Hemolysing solution at room temperature (15 to 30 °C) or refrigerated (2 to 8 °C). It is stable until the expiration date indicated on the kit package or Hemolysing solution vial label. DO NOT FREEZE.

Once the Hemolysing solution vial has been opened and positioned on the CAPILLARYS 3 instrument, it is stable for a maximum of 3 months (accumulated). If the Hemolysing solution vial is planned to be used for more than 3 months, it must be removed from the instrument after each use and stored at room temperature (15 to 30 °C) or refrigerated (2 and 8 °C), Hemolysing solution is then stable until the expiration date indicated on the Hemolysing solution vial label.

Discard Hemolysing solution if it changes its appearance, e.g., becomes cloudy due to microbial contamination.

3. FILTERS

Use

Disposable filters for filtration of analysis buffer, hemolysing solution and distilled or deionized water (used for capillaries rinsing).

IMPORTANT : When kit replacement, change systematically all the filters. Wear clean gloves for handling and installation of filters.

Screw one filter at the connector situated at the extremity of each tube that plunges in the vials of buffer, hemolysing solution and distilled or deionized water. When setting filters on the instrument, rinse the connectors and the tubes with distilled or deionized water.

Storage

Before use, store the filters in their sealed package in a dry place at room temperature (15 to 30 °C) or refrigerated (2 to 8 °C).

REAGENTS REQUIRED BUT NOT SUPPLIED WITH THE KIT

WARNING : *See the safety data sheets.*

1. NORMAL Hb A2 CONTROL

Intended use

The Normal Hb A2 Control (SEBIA, PN 4778) is designed for the migration control and for the quality control of human hemoglobin A2 quantification with CAPI 3 HEMOGLOBIN(E) electrophoresis procedure performed with the CAPILLARYS 3 automated instrument for capillary electrophoresis. The values obtained must fall within the range provided with each batch of Normal Hb A2 Control.

Composition

The Normal Hb A2 Control is obtained from a pool of normal human blood samples. The Normal Hb A2 Control is in a stabilized lyophilised form.

Use

IMPORTANT : For optimal use of the Normal Hb A2 Control with the CAPILLARYS 3 instrument, it is necessary to use one specific tube designed for blood controls and its corresponding cap (see "EQUIPMENT AND ACCESSORIES REQUIRED", Tubes and caps for Controls) and to identify this tube with the Normal Hb A2 Control bar code label.

- Reconstitute each lyophilized Normal Hb A2 Control vial with the volume of distilled or deionized water indicated in the instructions for use of the Normal Hb A2 Control. Allow to stand for 30 minutes and mix gently (avoid formation of foam).
NOTE : The precision of the reconstitution volume to be maintained is ± 1.0 %.
- Prepare 2 aliquots with equivalent volumes (≈ 0.850 mL) of the whole amount of the reconstituted control in conical tubes for control blood and close the tubes with their caps.
- Identify each tube with a Normal Hb A2 Control bar code label.

Migration control :

For the migration control, the recommendations to analyze the Normal Hb A2 Control are the following :

- Perform 1 series of analyses with the control :
 - before starting a new analysis sequence,
 - at the end of an analysis sequence.
 - Perform 2 successive series of analyses with the control :
 - after having changed the lot number of analysis buffer,
 - after having changed the technique,
 - after a capillary cleaning sequence with CAPICLEAN,
 - after a software upgrade,
 - after capillaries activation.
 - Perform 3 successive series of analyses with the control :
 - for the first use of the "HEMOGLOBIN(E)" analysis program with the CAPILLARYS 3 instrument,
 - after a prolonged stoppage (over 1 week).
- Place a tube with the reconstituted Normal Hb A2 Control in position No. 1 on the CAPILLARYS 3 sample rack No. 0 (store the second tube according to the indications of the Normal Hb A2 Control instructions for use).
 - Slide the sample rack No. 0 into the CAPILLARYS 3 instrument, the analysis starts automatically.
 - In the window which appears on the screen, select the number of analyses of the control to perform and validate.
 - The results are then automatically considered by the software for the data analysis.
On the review window and on the profile displayed in mosaic format, the symbol "a" indicates that the analysis of the migration control has been performed with an automatic dilution. The symbol "r" indicates that the analysis has been performed by successive re-injections of the diluted control contained in the reagent cup that has previously been analyzed (according to the number of analyses selected by the operator).

IMPORTANT : The hemoglobin A fraction of the Normal Hb A2 Control must show a minimal optical density (OD) of 0.10. Under this value, the recentering of the electrophoretic pattern will not occur correctly. When analysing samples, the identification of hemoglobin fractions, Hb A, Hb F, Hb A2 and Hb C and also the determination of the migration zone of other variants, may be impossible or wrong (see the paragraph RESULT ANALYSIS).

NOTE : After the installation of CAPILLARYS 3 instrument, during the first sequence of blood sample analysis, a red warning signal will appear if hemoglobin A is absent in one sample (and the recentering of the electrophoretic pattern will not be possible, see paragraph "Result analysis").

It is then recommended to analyze a blood sample with hemoglobin A on the concerned capillary and to analyze again the sample without hemoglobin A by placing it in a position corresponding to a capillary which has already detected hemoglobin A.

Quality control :

It is recommended to include one analysis of Normal Hb A2 Control into each run of samples, it should be used as a normal human blood. After reconstitution, analyze directly one of the aliquots of the Normal Hb A2 Control (applied in a tube for control with its cap and identified with one bar code label) as a blood sample to analyze on a sample rack. It will be automatically diluted with hemolysing solution. The Normal Hb A2 Control may also be analyzed with the sample rack No. 0, see paragraph before (Migration control). The values obtained must fall within the range provided with each batch of Normal Hb A2 Control.

Storage, stability and signs of deterioration

See the Normal Hb A2 Control instructions for use.

NOTE : For optimal use with the CAPILLARYS 3 instrument, it is recommended to prepare 2 aliquots with equivalent volumes (≈ 0.850 mL) in conical tubes for controls of the reconstituted Normal Hb A2 Control before freezing it.

WARNING : No test method can provide an absolute assurance of the absence of HIV, hepatitis B and C or other infectious agents. Therefore, handle the Normal Hb A2 Control as a hazardous biological material.

This lot of control blood was found negative on assays approved by FDA or EU equivalent regulatory agency :

- against hepatitis B surface antigen,
- for antibody to HCV,
- for antibody to HIV1 and HIV2.

2. DISTILLED OR DEIONIZED WATER

Use

For capillaries rinsing in automated instrument for capillary electrophoresis CAPILLARYS 3, SEBIA.

It is recommended to use filtered distilled or deionized water (on a filter with a porosity ≤ 0.45 μm) and with a conductivity lower than 3 $\mu\text{S/cm}$, which corresponds to a resistivity higher than 0.33 M Ω .cm.

To prevent microbial proliferation, change the water every day.

For optimal operation, add CLEAN PROTECT (SEBIA, PN 2059, 1 vial of 5 mL) in distilled or deionized water (see the instructions for use of CLEAN PROTECT) or use directly the ready to use CAP|protect[®] solution (SEBIA, PN 2061 : 2 containers of 5 L of distilled water with CLEAN PROTECT).

IMPORTANT : Before filling the rinse container, it is recommended to wash it with plenty of distilled or deionized water.

** NOTE : The CAP|protect solution can also be used to dilute the stock wash solution. Then, in that case, the diluted wash solution may show a transient more or less marked yellow colour without any adverse effects on its performance.*

3. CAPILLARYS 3 CAPICLEAN

Composition

The vial of CAPICLEAN concentrated solution (SEBIA, PN 2060, 1 vial of 25 mL) contains : proteolytic enzymes, surfactants and additives nonhazardous at concentrations used, necessary for optimum performances.

Use

For sample probe cleaning in automated instrument for capillary electrophoresis CAPILLARYS 3, SEBIA, during the CAPICLEAN cleaning sequence.

IMPORTANT :

- When less than 500 samples are analyzed within a week, launch a CAPICLEAN cleaning sequence at least once a week.
- When less than 500 samples are analyzed within a day but more than 500 samples are analyzed within a week, launch a CAPICLEAN cleaning sequence after every 500 analyses.
- When more than 500 samples are analyzed within a day, launch a CAPICLEAN cleaning sequence once a day.

See the instruction for use of CAPILLARYS 3 CAPICLEAN and the instruction manual of CAPILLARYS 3, SEBIA.

Storage, stability and signs of deterioration

See the instructions for use of CAPILLARYS 3 CAPICLEAN, SEBIA.

4. SODIUM HYPOCHLORITE SOLUTION (for sample probe cleaning)

Preparation

Prepare a sodium hypochlorite solution (2 % to 3 % chloride) by diluting 250 mL 9.6 % chloride concentrated solution to 1 liter with cold distilled or deionized water.

Use

For the sample probe cleaning in the CAPILLARYS 3 instrument, SEBIA (weekly maintenance in order to eliminate adsorbed proteins from the probe).

See the CAPILLARYS 3 instruction manual, SEBIA.

Storage, stability and signs of deterioration

Store the working chlorinated solution at room temperature (15 to 30 °C) in a closed container, it is stable for 3 months. Avoid storage in sunlight, close to heat and ignition source, and to acids and ammonia.

5. CAPILLARYS 3 WASH SOLUTION

Preparation

The vial of the stock wash solution (SEBIA, PN 2062, 1 vial, 75 mL) should be diluted up to 750 mL with distilled or deionized water.

After dilution, the wash solution contains an alkaline solution pH ≈ 12.

Use

For washing the capillaries before electrophoretic separation.

IMPORTANT :

- When wash solution vial replacement, change systematically the filter. Wear clean gloves for handling and installation of the filter.
- Before placing the wash solution vial in the instrument, it is recommended to wash the connector and the tube with plenty of distilled or deionized water to avoid salts deposit.
- Screw the filter at the connector situated at the extremity of the tube plunging in the wash solution vial.

Storage, stability and signs of deterioration

Store the stock and working wash solutions in closed containers at room temperature (15 to 30 °C) or refrigerated (2 and 8 °C).

The stock wash solution is stable until the expiration date indicated on the kit or wash solution vial label.

Working wash solution is stable for 3 months.

After dilution and immediate installation of the vial in the instrument, the solution is stable for 3 months (if the working wash solution is stored out of the instrument before use, this time of 3 month storage must take into account the time during which the solution is stored outside the instrument).

Discard working wash solution if it changes its appearance, e.g., becomes cloudy due to microbial contamination.

6. SALINE

Preparation

Make 0.15 M (0.9 g/dL) NaCl solution in distilled or deionized water.

Use

To analyze samples with an additional fraction in Z(C) migration zone (Hb C migration zone) or Z(A2) migration zone (Hb A2 migration zone) (see § Sample preparation, Particular cases).

Storage, stability and signs of deterioration

Store saline at room temperature (15 to 30 °C) or refrigerated (2 - 8 °C).

Discard after 3 months or if it changes its appearance, e.g., becomes cloudy due to microbial contamination. For longer storage periods, add sodium azide, 0.1 g/dL.

OPTIONAL REAGENT BUT NOT SUPPLIED

WARNING: See the safety data sheet.

PATHOLOGICAL Hb A2 CONTROL

The Pathological Hb A2 Control, SEBIA, PN 4779, can be used for the migration control, in addition or as a replacement of the Normal Hb A2 Control. For its utilization for the migration control or quality control, the Pathological Hb A2 Control should be used like the Normal Hb A2 Control, see the previous paragraph "NORMAL Hb A2 CONTROL".

See the instructions for use of the Pathological Hb A2 Control for additional information.

NOTES :

The assays that were performed for the validation of reagents demonstrated that, for the different solutions and using an adapted equipment for the reconstitution volume, a variation of $\pm 5\%$ on the final volume has no adverse effect on the analysis.

The distilled or deionized water used to reconstitute solutions, must be free of bacterial proliferation and mold (use a filter $\leq 0.45 \mu\text{m}$) and have a conductivity lower than $3 \mu\text{S/cm}$, which corresponds to a resistivity higher than $0.33 \text{M}\Omega\cdot\text{cm}$.

EQUIPMENT AND ACCESSORIES REQUIRED NOT INCLUDED IN THE KIT

1. SEBIA CAPILLARYS 3 instrument for capillary electrophoresis : CAPILLARYS 3 OCTA PN 1245 or CAPILLARYS 3 TERA PN 1246, connected to a computer equipped with the PHORESIS software for data processing.
2. Sample racks supplied with CAPILLARYS 3 instrument.
3. CAPILLARYS 3 & MC SWITCH RACK FOR HEMOGLOBIN(E) (1), SEBIA, PN 1373, to launch automatically a technique change to HEMOGLOBIN(E) procedure on the CAPILLARYS 3 instrument.
4. CAPILLARYS 3 & MC LOW VOLUMES RACKS (5), SEBIA, PN 1364, for the analysis of samples with volume below $800 \mu\text{L}$ on the CAPILLARYS 3 instrument.
5. Container kit supplied with CAPILLARYS 3 instrument : Rinse (to fill with distilled or deionized water), wash solution and waste container.
6. CAPI 3 REAGENT CUPS (24 x 14), SEBIA, PN 2582, including 24 packs of 14 CAPI 3 reagent cups : Single use cups for the preparation of biological samples to analyze with the automated instrument. To be placed on the automated loading system for cups of CAPILLARYS 3. One reagent cup is intended for the analysis of 8 samples with CAPILLARYS 3 OCTA and 12 samples with CAPILLARYS 3 TERA.

WARNING : After use, reagent cups with biological samples have to be handled with care. When the analysis is completed, reagent cups must be discarded with biological waste products and they must NEVER be reused.

Storage : Before use, store the reagent cups in their sealed package in a clean and dry place and at a temperature comprised between 2 and 30°C .

7. CAPI 3 BINS FOR USED REAGENT CUPS (5), SEBIA, PN 2581 : Bins intended for automated collection of used reagent cups in CAPILLARYS 3. To place in CAPILLARYS 3 at the location intended for this purpose.

WARNING : Bins containing used reagent cups with biological samples have to be handled with care.

8. Collection tubes with 13 mm diameter and their corresponding caps (maximal length of tube with cap : 91 mm, maximal diameter of cap : 17 mm) : for example, BD Vacutainer, Terumo Venosafe 5 mL, Greiner Bio-one Vacuette 1, 2, 3 or 4 mL or Sarstedt S-Monovette 2.6, 2.7 or 3.4 mL tubes (13 x 75 mm),

or

collection tubes with 11 mm diameter and their corresponding caps (maximal length of tube with cap : 91 mm, maximal diameter of cap : 17 mm) : for example, Sarstedt S-Monovette 2.7 mL or Kabe Labortechnik Primavette S 2.6 mL tubes (11 x 66 mm),

or collection tubes with equivalent dimensions approved for clinical assays.

WARNING :

- Do not use these collection tubes on a sample rack No. 0 from the CAPILLARYS 3 instrument (the sample rack No. 0 must only be used with conical tubes for the analysis of blood controls).
- Do not use Sarstedt S-Monovette collection tubes on a CAPILLARYS 3 & MC LOW VOLUMES rack (important risk of instrument and tube damage).

9. TUBES AND CAPS FOR CONTROLS, SEBIA, PN 9202 (20 units) or PN 9205 (500 units) : conical tubes and their caps to analyze blood controls and samples with a low volume (see § Sample preparation), with the CAPILLARYS 3 instrument.
10. TEST TUBES, SEBIA, PN 9214 : 200 100 mm-tubes for the hypochlorite sodium solution intended for the cleaning of the sample probe, or tubes (without cap) with equivalent dimensions (length comprised between 90 and 100 mm and diameter comprised between 13 and 16 mm).

SAMPLES FOR ANALYSIS**Sample collection and storage**

Fresh anticoagulated whole blood samples collected in tubes containing KEDTA as anticoagulant are recommended for analysis. Blood must be collected according to established procedures used in clinical laboratory testing.

Samples can be stored for 7 days maximum between 2 and 8°C or 24 hours maximum at room temperature (between 15 and 30°C).

Progressive hemoglobins (Hb) degradation may occur for samples stored between 2 to 8°C .

When the blood sample is stored for more than 7 days at $2 - 8^\circ\text{C}$:

- a weak fraction, corresponding to methemoglobin, appears in the Hb S migration zone,
- when Hb C is present, a fraction corresponding to degraded Hb C appears more anodic than Hb A2 which does not interfere with it (Z(E) zone),
- when Hb O-Arab is present, a fraction corresponding to degraded Hb O-Arab appears in the Hb S migration zone (Z(S) zone),
- when Hb E is present, a fraction corresponding to degraded Hb E appears in the Z(D) zone,
- when Hb S is present, a fraction corresponding to degraded Hb S appears in the Hb F migration zone (Z(F) zone),
- when Hb A is present, a fraction corresponding to degraded Hb A ("aging fraction" of Hb A) appears more anodic (Z11 zone).

When Hb F is present (in blood samples from newborn babies), a fraction appears in the Hb A migration zone (Z(A) zone) due to the sample degradation.

When stored for more than 10 days, viscous aggregates in red blood cells are observed ; it is necessary to discard them before the analysis.

For longer storage, whole blood samples should be frozen quickly at $-70 / -80^\circ\text{C}$ (within 8 hours maximum after collection) without prior preparation. Frozen whole blood samples are stable for 3 months maximum at $-70 / -80^\circ\text{C}$.

IMPORTANT : For optimal storage of blood samples, do not store them at -20°C but at -80°C (see BIBLIOGRAPHY, J. Bardakjian-Michau *et al*, 2003).

Sample preparation

- Use directly whole blood samples.
- Check that all the tubes contain 800 µL minimum of blood and are perfectly closed.
- **Vortex for 5 seconds blood samples stored at 2 - 8 °C for one week or stored at - 70 / - 80 °C.**

WARNING : The tubes must be closed with their corresponding caps designed for the CAPI 3 HEMOGLOBIN(E) procedure performed with the CAPILLARYS 3 instrument (see EQUIPMENT AND ACCESSORIES REQUIRED).

Particular cases:

Analysis of samples without any Hb A and with Hb F < 3 % or without any Hb A2 (these samples are perfectly quantified but not identified by zones). To identify hemoglobin fractions in a sample without any Hb A and with Hb F < 3 % or without any Hb A2, it is recommended to prepare this sample according to the following procedure:

- Vortex for 5 seconds the whole blood sample.
- In a conical tube for control, mix one volume (50 µL) of whole blood to analyze with one volume (50 µL) of Normal Hb A2 Control and cap the tube.
- Vortex for 5 seconds.
- Place the tube on a CAPILLARYS 3 & MC LOW VOLUMES rack.
- Slide the rack into the CAPILLARYS 3 instrument.
- Perform the analysis of this sample according to the standard procedure like a usual blood sample.

The results are then automatically considered by the software for the data analysis.

IMPORTANT : For a sample without any Hb A, Hb F or Hb A2 prepared according to this procedure, the result obtained with the mixed sample will enable presumptive variant identification due to the positioning of the hemoglobins fractions in the appropriate identification zones. Do not report the relative quantification from the mixed sample result.

The relative quantification of hemoglobins should be reported utilizing the initial, unmixed sample result (without any dilution in the blood control).

Analysis of a sample with an additional fraction in Z(C) migration zone (Hb C migration zone) or Z(A2) migration zone (Hb A2 migration zone) :

The presence of a Hb Constant Spring variant may be suspected when a hemoglobin fraction is observed in Z(C) or Z(A2) migration zones. This fraction may also be due to plasmatic proteins from the sample (from a patient with anaemia for example, with a decreased [red blood cells] / [plasma] ratio).

The analysis of red blood cells from the same sample, without plasmatic proteins, will confirm the presence of this variant.

Prepare the sample according to the following procedure :

- Centrifuge the whole blood sample to obtain a red blood cells pellet, discard plasma.
- In a conical tube for control, mix one volume (50 µL) of red blood cells with one volume (50 µL) of saline and cap the tube.
- Vortex for 5 seconds.
- Place the tube on a CAPILLARYS 3 & MC LOW VOLUMES rack.
- Slide the rack into the CAPILLARYS 3 instrument.
- Perform the analysis of this sample according to the standard procedure like a usual blood sample.

The results are then automatically considered by the software for the data analysis.

Analysis of samples with a low volume

The following table presents the tubes and sample racks to use according to the minimum volume of sample to analyze.

	STANDARD TUBE	STANDARD TUBE (EXCEPT FOR Sarstedt S-Monovette tubes)	TUBES AND CAPS FOR CONTROLS (conical tubes, PN 9202 & 9205)	
	CAPILLARYS 3 & MC SAMPLE RACKS (PN 1369)	CAPILLARYS 3 & MC LOW VOLUMES RACKS (PN 1364)	CAPILLARYS 3 & MC SAMPLE RACKS (PN 1369)	CAPILLARYS 3 & MC LOW VOLUMES RACKS (PN 1364)
Minimum volume of sample needed for the CAPI 3 HEMOGLOBIN(E) analysis	800 µL	300 µL (1)	400 µL (2)	100 µL (3)
Software version for HEMOGLOBIN(E)	≥ 1.08	≥ 1.08	≥ 1.08	≥ 1.08
Handling	No handling of the sample --> complete traceability	No handling of the sample --> complete traceability	Apply a minimum of 400 µL of sample in a conical tube	Apply a minimum of 100 µL of sample in a conical tube

(1) Analysis of samples with a volume comprised between 300 and 800 µL (EXCEPT for Sarstedt S-Monovette tubes) :

- Place the capped tube with whole blood to analyze (at least 300 µL) on a CAPILLARYS 3 & MC LOW VOLUMES rack.
- Slide the rack into the CAPILLARYS 3 instrument.

WARNING : Do not use Sarstedt S-Monovette collection tubes on a CAPILLARYS 3 & MC LOW VOLUMES rack (important risk of instrument and tube damage). For samples with a volume below 800 µL in this type of tube, follow the procedure that corresponds to the volume to analyze.

(2) Analysis of samples with a volume comprised between 400 and 800 µL (for Sarstedt S-Monovette tubes in particular) :

- Vortex for 5 seconds the whole blood sample to analyze.
- Apply in a conical tube for control the whole blood sample (at least 400 µL) and cap the tube.
- Identify the tube with the specific bar code label of the sample.
- Place the tube on a CAPILLARYS 3 & MC SAMPLE rack.
- Slide the rack into the CAPILLARYS 3 instrument at the beginning of an analysis series.

NOTE : It is recommended to gather samples with volume comprised between 400 and 800 µL on the same sample rack and analyze them at the beginning of an analysis series. Mix well the sample applied in a conical tube for the analysis before sliding the sample rack into the automated instrument. Without any bar code label on the conical tube, the sample cannot be identified.

(3) Analysis of samples with a volume comprised between 100 and 300 µL :

- Vortex for 5 seconds the whole blood sample to analyze.
- Apply in a conical tube for control the whole blood sample (at least 100 µL) and cap the tube.
- Identify the tube with the specific bar code label of the sample.
- Place the tube on a CAPILLARYS 3 & MC LOW VOLUMES rack.
- Slide the rack into the CAPILLARYS 3 instrument at the beginning of an analysis series.

NOTE : It is recommended to gather samples with volume comprised between 100 and 300 µL on the same sample rack and analyze them at the beginning of an analysis series. Mix well the sample applied in a conical tube for the analysis before sliding the rack into the automated instrument. Without any bar code label on the conical tube, the sample cannot be identified.

Samples to avoid

- Avoid coagulated blood samples.
 - Avoid aged, improperly stored blood samples ; the automated hemolysis of samples may be disturbed by viscous aggregates in red blood cells. Then, degradation products (as artefacts) may affect the electrophoretic pattern.
- In these 2 previous cases, aggregates in red blood cells may affect the collection of the sample by the probe.
- Do not analyze directly tubes containing less than 800 µL of blood sample, the analysis should be affected (see particular cases).
 - Do not use samples from neonate / newborn population. The CAPI 3 HEMOGLOBIN(E) procedure performed with the CAPILLARYS 3 instrument has not been evaluated in the neonate / newborn population (age range – birth to 28 days).

PROCEDURE

The CAPILLARYS 3 instrument is a multiparameter instrument for hemoglobins analysis on parallel capillaries. The hemoglobins assay uses 8 or 12 capillaries to run the samples.

The sequence of automated steps is as follows :

- sample racks identification by RFID (Radio Frequency Identification),
- bar code reading of sample tubes (for up to 8 tubes),
- mixing of blood samples before analysis,
- sample hemolysis and dilution from primary tubes into reagent cups,
- capillary washing,
- injection of hemolyzed samples,
- hemoglobin separation and direct detection of the separated hemoglobins on capillaries.

The manual steps include :

- placement of reagents and disposables into the CAPILLARYS 3 instrument,
- placement of sample tubes (with caps) in sample racks,
- placement of racks on the CAPILLARYS 3 instrument,
- removal of sample racks and sample tubes after analysis,
- removal of bins for used reagent cups.

PLEASE CAREFULLY READ THE CAPILLARYS 3 INSTRUCTION MANUAL.

I. PREPARATION OF ELECTROPHORETIC ANALYSIS

1. Switch on CAPILLARYS 3 instrument and computer.
2. Wait until the instrument is completely initialized.
3. Start the PHORESIS software installed on the computer for data processing.
4. The CAPI 3 HEMOGLOBIN(E) kit is intended to run with "HEMOGLOBIN(E)" analysis program from the CAPILLARYS 3 instrument. To select "HEMOGLOBIN(E)" analysis program and place the CAPILLARYS HEMOGLOBIN(E) buffer and hemolyzing solution vials in the instrument, please read carefully the CAPILLARYS 3 instruction manual. If necessary, place the vial with the reconstituted wash solution in the instrument.
5. The sample rack contains 8 positions for sample tubes. Place up to 8 capped sample tubes with whole blood on each sample rack ; the bar code of each tube must be visible in the openings of the sample rack.
6. Take a pack of new reagent cups by holding the handle and place it on the automated loading system for cups of CAPILLARYS 3 ; then, remove the flange (a message will be displayed when reagent cups are missing).
7. Place a new bin for used reagent cups into the CAPILLARYS 3 instrument at the location intended for this purpose.
8. Slide the sample rack(s) into the CAPILLARYS 3 instrument through the opening in the right side of the instrument. Up to 15 sample racks can be introduced successively and continuously into the instrument.

NOTES :

- When analyzing a control blood sample, it is advised to use specific conical tubes for control bloods and their corresponding caps, and a rack No. 0 for controls or a sample rack.
- Do not analyze blood samples on a sample rack No. 0, the analysis should be affected.

9. Remove analyzed sample racks from the plate on the left side of the instrument.
10. If necessary, take off carefully the bin containing used reagent cups and discard it.

WARNING : Bins containing used reagent cups with biological samples have to be handled with care.

DILUTION - MIGRATION - DESCRIPTION OF THE AUTOMATED STEPS

1. Sample rack identification by RFID.
2. Bar codes are read on primary sample tubes.
3. Mixing of tubes.
4. Samples are diluted in hemolysing solution and the sample probe is rinsed after each sample.
5. Capillaries are washed.
6. Diluted samples are injected into capillaries.
7. Migration is carried out under constant voltage for about 8 minutes and the temperature is controlled by Peltier effect.
8. Hemoglobins are detected directly by scanning at 415 nm and data of the obtained hemoglobin electrophoretic pattern are transmitted from the instrument to the computer equipped with the software for data processing.

II. RESULT ANALYSIS

At the end of the analysis, the corresponding data are transmitted by the instrument to the software for data processing and a hemoglobin electrophoretic profile appears on the screen of the computer. Relative quantification of individual hemoglobin fractions is automatically performed and profiles can be analyzed. The hemoglobin fractions Hb A, Hb F, Hb A2 and Hb C are automatically identified. The Hb A fraction is centered in the middle of the review window and Hb A2 is adjusted at a fixed position against that of Hb A. In the absence of Hb A and when Hb F is present ($\geq 3\%$), the recentering of the pattern is made with Hb F and Hb A2 peaks that are placed at fixed positions.

The resulting electrophoregrams are evaluated visually for pattern abnormalities.

The electrophoretic patterns are colored :

- in Cyan when the number of fractions / peaks is that which is configured by default for the procedure (2 fractions for HEMOGLOBIN(E) procedure, for example),
- in Magenta when the number of fractions / peaks is not that which is configured by default for the procedure.

With HEMOGLOBIN(E) procedure, the Hb F peak is orange (identified by « Hb F or variant ») when the age of the patient is unknown and blue (identified by « Hb F ») when the age of the patient is known and the fraction / peak is lower than 2%.

The resulting electrophoregrams are evaluated visually for pattern abnormalities.

The potential positions of the different hemoglobin variants (identified in zones called Z1 to Z15) are shown on the screen of the system and indicated on the result ticket.

See the table with known variants which may be present in each corresponding zone.

When the software identifies a hemoglobin fraction in a defined zone, the name of this zone is framed.

Patterns are automatically adjusted with regard to Hb A and Hb A2 fractions, or with regard to Hb F and Hb A2 fractions as the case may be, to facilitate their interpretation:

- when Hb A and / or Hb A2 fractions are not detected on an electrophoretic pattern and / or when Hb F (with no Hb A) is not detected or is at a level $< 3\%$,
 - a yellow warning signal appears,
 - the adjustment of the pattern is performed using the position of the Hb A fraction on the two previous patterns obtained with the same capillary,
 - no fraction is identified (except when Hb C is detected: in this case, Hb A2 and Hb C fractions are identified),
 - the different migration zones (Z1 to Z15) do not appear neither on the screen of the system, nor on the ticket result.
- when Hb F is detected at a level $\geq 3.0\%$, without any detection of Hb A (no Hb A or Hb A at a low level) on an electrophoretic pattern,
 - the adjustment of the pattern is performed using the position of the Hb F and Hb A2 fractions,
 - Hb F and Hb A2 fractions are placed at fixed positions,
 - Hb F and Hb A2 fractions are identified,
 - the different migration zones (Z1 to Z15) are indicated on the screen of the system and on the ticket result by the same way of patterns with Hb A,
 - abnormal fractions are grey-dashed and identified using their migration zone (a fraction detected in Z(D) zone is called "Z(D) zone" for example),
 - when a rare variant migrates in the Hb A2 migration zone, the different migration zones (Z1 to Z15) do not appear neither on the screen of the system, nor on the ticket result.
- when the adjustment is not possible,
 - a red warning signal appears,
 - no fraction is identified,
 - the different migration zones (Z1 to Z15) do not appear neither on the screen of the system, nor on the ticket result.

Call SEBIA.
- when optical density (OD) is insufficient on a migration control electrophoretic pattern (obtained with the Normal Hb A2 Control or the Pathological Hb A2 Control, identified with its bar code label on the sample rack No. 0),
 - a warning message is displayed in order to consider or remove this analysis for the determination of Hb A fraction position,
 - a purple warning signal appears on the review window,
 - Hb A and Hb A2 fractions are not identified (except when the analysis is considered by the operator),
 - the different migration zones (Z1 to Z15) do not appear neither on the screen of the system, nor on the ticket result (except when the analysis is considered by the operator).

In all cases, the different migration zones (Z1 to Z15) do not appear neither on the screen of the system, nor on the ticket result.

On the electrophoretic pattern, the curves of Hb A2 and Hb C fractions, are calculated and redrawn by fitting with adjustment (or fitted) and are overlaid with the native curve. This display allows the Hb A2 fraction quantification if Hb C is present in the sample.

WARNING : In some cases of hemoglobin C (homozygous) or after a technical problem, the hemoglobins A2 and C are not fitted ; these fractions are then under-quantified. It is then recommended to quantify the Hb A2 fraction by using another technique.

PLEASE CAREFULLY READ THE PHORESIS INSTRUCTION MANUAL.

III. END OF ANALYSIS SEQUENCE

At the end of each analysis sequence, the operator must initiate the "shut down" procedure of the CAPILLARYS 3 instrument in order to store capillaries in optimal conditions.

IV. FILLING OF REAGENT CONTAINERS AND MANAGEMENT OF DISPOSABLES

The CAPILLARYS 3 instrument has an automatic control for reagents (by using RFID labels) and for disposables (reagent cups and bins for used cups).

IMPORTANT : It is necessary to respect the designed position for wash solution, rinse and waste containers.

On the screen of the CAPILLARYS 3 instrument, the "Main compartment" menu for reagents management displays information when it is necessary to perform one of the following tasks :

- place a new buffer vial and / or,
- place a new hemolysing solution vial and / or,
- place a new vial with working wash solution and / or,
- fill the container with filtered distilled or deionized water for rinsing capillaries and / or,
- empty the waste container.

WARNING : *Do not use marketed deionized water, such as water for ironing for example (risk of important capillaries damage). Use only water with ultrapure quality, such as injection grade water.*

IMPORTANT : Before filling the rinse container, it is recommended to wash it with plenty of distilled or deionized water.

PLEASE CAREFULLY READ THE CAPILLARYS 3 INSTRUCTION MANUAL.

QUALITY CONTROL

It is advised to include into each run of samples, an assayed control blood (for example, a blood sample containing hemoglobins A, F, C and S, such as Hb AFSC Control, SEBIA, PN 4792, or a normal blood sample, the Normal Hb A2 Control, SEBIA, PN 4778 or the Pathological Hb A2 Control, SEBIA, PN 4779).

IMPORTANT : For optimal use of the blood controls analyzed with the CAPILLARYS 3 instrument, it is necessary to use the specific conical tubes for controls and their corresponding caps (see "EQUIPMENT AND ACCESSORIES REQUIRED") and the bar code labels intended to identify the tubes for controls that contain the blood control to analyze.

RESULTS

Values

Direct detection at 415 nm in capillaries yields relative concentrations (percentages) of individual hemoglobin zones.

Reference values for individual major electrophoretic hemoglobin zones have been established from a healthy population of 113 adults (men and women) with normal hemoglobin values using HPLC technique :

Hemoglobin A : comprised between 96.7 and 97.8 %

Hemoglobin F : ≤ 0.5 % (*)

Hemoglobin A2 : comprised between 2.2 and 3.2 %

(*) See *Interference and limitations*

It is recommended that each laboratory establish its own threshold values.

NOTE : *Reference values have been established using the standard parameters of the software (smoothing 0 and hemoglobin fractions automatic quantification with HEMOGLOBIN(E) analysis program).*

WARNING : *Reference values must be considered only when hemoglobin variants are absent.*

Interpretation

See *ELECTROPHORETIC PATTERNS*, figures 1 – 18.

The different migration zones of hemoglobin variants (called Z1 to Z15) are shown on the screen of the system and on the result ticket. Passing the mouse cursor over a zone name displays icon information containing possible hemoglobin variants that could be seen in this zone.

For each fraction, the maximum position defines the migration zone.

See the table showing the potential variants located in each zone.

With PHORESIS VS ≥ 9.15 , this table lists 525 different hemoglobin variants. Due to the history of their discovery, some variants may have many names. A second name is added between brackets close to the main name (for example, in zone Z(D), Hb Korle-Bu (G-Accra)). Other names are not listed in this table.

In zone Z(A), variants are listed in alphabetical order.

For other zones, variants are sorted in main fractions and minor fractions and presented by migration order from most cathodic variants to most anodic variants.

For variants with a main fraction that migrates in zone Z(A), their minor fractions which migrate in zone Z(A2) are not indicated.

- The symbol "*" indicates a hidden or partially hidden peak due to similar migration to normal Hb A or Hb A2 fraction. A partially hidden fraction corresponds to a more or less important shoulder of the normal fraction.

- The symbol “#” indicates the display in icon information of several visible fractions from the same variant, generally present in different zones (for example, alpha-chain variant with a second visible peak as Hb Q-India, or unstable variant as Hb Sabine and Hb Köln). Not concerned: beta chain variants except unstable variants, gamma and delta chain variants and delta-beta hybrids, alpha chain variants without second peak visible on the electrophoretic pattern.
- The symbol “!” alerts the potential risk of a migration zone shift for a rare variant located in a zone boundary. Additionally, the migration variation of a variant (± 1 point) depends on its percentage. For example, Hb Willamette, located on the far right of zone Z(F), may migrate in zone Z(D) when its percentage has decreased in case of an associated thalassemia.

These symbols are explained in the “Captions” icon information located in the upper left side of the review window.

1. Qualitative abnormalities : Hemoglobinopathies

Most hemoglobinopathies are due to substitution by mutation of a single amino acid in one of the four types of polypeptide chains (1, 2, 4, 9, 12). The clinical significance of such a change depends on the type of amino acid and the site involved (13). In clinically significant disease, either the α -chain or the β -chain is affected.

More than 1400 variants of adult hemoglobin have been described (6, 14). The first abnormal hemoglobins studied and the most frequently occurring have an altered net electric charge, leading to an easy detection by electrophoresis.

There are five main abnormal hemoglobins which present a particular clinical interest : S, C, E, O-Arab and D.

The CAPI 3 HEMOGLOBIN(E) kit is intended for the identification of hemoglobinopathies and thalassemias.

Hemoglobin S

Hemoglobin S is the most frequent. It is due to the replacement of one glutamic acid (an acidic amino acid No. 6) of the β -chain by valine (a neutral amino acid) : when compared to Hb A, its isoelectric point is elevated and its total negative charge decreased with the analysis pH. Its electrophoretic mobility is therefore increased in the capillary and this hemoglobin is faster than A fraction. With alkaline buffered CAPI 3 HEMOGLOBIN(E) procedure, hemoglobin S migrates between A and A2 fractions, next to Hb A2.

Hemoglobin C

One glutamic acid of the β -chain is replaced by lysine (a basic amino acid No. 6) : its mobility is strongly reduced. When compared to Hb A, its isoelectric point is highly elevated and its total negative charge decreased with the analysis pH. Its electrophoretic mobility is therefore increased in the capillary and this hemoglobin is faster than A fraction which allows its differentiation. Hemoglobins C, E and O-Arab are not superimposed on the electrophoretic pattern and are easily identified.

Hemoglobin E

One glutamic acid of the β -chain (No. 26) is replaced by lysine. With CAPI 3 HEMOGLOBIN(E) procedure, hemoglobin E migrates just anodically behind hemoglobin A2 and is totally separated from it. Then, when hemoglobin E is present, A2 fraction can be measured to detect β -thalassemia.

Hemoglobin O-Arab

One glutamic acid of the β -chain (No. 121) is replaced by lysine. With CAPI 3 HEMOGLOBIN(E) procedure, hemoglobin O-Arab migrates exactly like hemoglobin A2. In such a case, hemoglobin A2 can not be quantified. When this fraction is $> 10.5\%$, hemoglobin O-Arab must be suspected.

Note that Hb O-Arab migrates separately from hemoglobins C and E.

Hemoglobin D (Los Angeles)

One glutamic acid of the β -chain (No. 121) is replaced by glutamine. With CAPI 3 HEMOGLOBIN(E) procedure, hemoglobin D (called D-Punjab, D-Los Angeles, D-Chicago or D-Portugal) migrates behind hemoglobin S, this property allows to differentiate S and D hemoglobins.

2. Quantitative abnormalities : Thalassemias

Thalassemias constitute a quite heterogeneous group of genetic disorders characterized by decreased synthesis of one type of the polypeptide chains. The molecular mechanism of this decrease has not been fully described.

There are two types of thalassemia syndromes :

Alpha-thalassemias

They are characterized by the decrease of synthesis of the α -chains, consequently affecting the synthesis of all normal hemoglobins. The excess of synthesis of the β - and γ -chains in relation to α -chains induces the formation of tetramers without any α -chain :

- hemoglobin Bart = γ_4 ,
- hemoglobin H = β_4 .

Hemoglobin H presents a low isoelectric point ; with CAPI 3 HEMOGLOBIN(E) procedure, it migrates more anodic than hemoglobin A (and may appear as one or several fractions).

Beta-thalassemias

They are characterized by the decrease of synthesis of the β -chains. Only hemoglobin A synthesis is affected.

Therefore hemoglobin F and hemoglobin A2 percentages are increased with respect to hemoglobin A. With CAPI 3 HEMOGLOBIN(E) procedure, values obtained for different normal hemoglobin fractions allow the detection of beta-thalassemias.

3. Particular cases

- When there is no hemoglobin A in the sample, a small fraction may be observed in anodic position compared with Hb F (in the Z β zone when migration zones are displayed on the electrophoretic pattern) ; this fraction may be acetylated hemoglobin F which represents about 15 to 25 % of hemoglobin F. The CAPILLARYS 3 system can identify this acetylated hemoglobin separately from the hemoglobin A without any confusion.
- When a small fraction (about 0.5 to 3 %) migrates between hemoglobins F and δA^2 (A2 variant), a hemoglobin A2 variant may be suspected.
- When a hemoglobin A2 variant is detected (δA^2 or any other A2 variant), it is recommended to add its percentage to hemoglobin A2 for a better beta-thalassemia diagnostic.
- Some hemoglobin variants (such as Hb Camperdown and Hb Okayama) migrate close to Hb A and may not be separated from this hemoglobin.
- Some hemoglobin variants (such as Hb Pórtó-Alegre or degraded Hb S, for example) including homozygous variants such as Hb Q-Thailand, migrate close to Hb F. In the absence of Hb A, the adjustment of the pattern using Hb F and Hb A2 peaks and the display of migration zones prevents any confusion of these variants with Hb F.
- In zone Z12, the curve of Hb Bart is calculated and redrawn by fitting with adjustment (or fitted). Fitted fractions are then called “Hb Bart suspected”. Narrow fractions with low percentage are not Hb Bart's, they are identified “Z12 zone”. Wide fractions with elevated percentage, suspected to be hemoglobin variants, are identified “Hb Bart zone”.

- In the CAPI 3 HEMOGLOBIN(E) technique, for diabetic patients with elevated HbA_{1c} (over 10 %), a small fraction is observed and eventually identified as a peak in Z10 zone.
- Weak fractions may be observed in Z14 and Z15 migration zones. It is then necessary to analyze the hematologic state of the patient and to perform complementary analyses in order to characterize these fractions (artefact or hemoglobin abnormality). **The software version (≥ 9.15) allows a specific identification of Hb H in Z15 zone. Fractions with a width over 10 points and a percentage between 0.3 and 32 % are called "Hb H suspected". Fractions with a width below 10 points are not Hb H and are identified "Z15 zone". Wide fractions with a percentage between 10 and 58 %, suspected to be hemoglobin variants, are identified "Abnormal Hb".**
- When analyzing blood samples from newborn babies, Hb A from samples containing Hb F at high concentrations may be disturbed, especially due to the presence of degraded Hb F in its migration zone. The Hb A percentage indicated by the software may be overvalued. In addition, when hemoglobin variants (> 4 %, such as Hb S, Hb C, Hb E or Hb D-Punjab) are present in blood samples containing high Hb F levels (> 60 %), it is necessary to perform complementary analyses in order to confirm the presence of Hb A.
- For newborn babies until 6 – 9 months old, it is recommended to analyze many blood samples (collected monthly, for example) in order to check the Hb F concentration. It will allow to verify the decrease of Hb F concentration and the potential presence of a variant. In case of uncertainty, it is advised to confirm by using complementary studies and to analyze parents' blood samples.

Examples with increased hemoglobin F (Hb F) (except for newborn babies) :

- pregnancy,
 - patients with sickle cell disease, more than 2 years old, with a Hydrea® (hydroxyurea) treatment and / or transfused and / or producing naturally Hb F increased by compensation,
 - patients, aged more than 2 years old, with HPPH trouble (hereditary persistence of foetal hemoglobin exhibiting 15 to 35 % Hb F for heterozygous patients),
 - patients, more than 2 years old, with leukaemia (with any type), hereditary haemolytic anemia, diabetes, thyroid disease, hyperactivity of bone marrow, multiple myeloma, cancer with metastases.
- Hb S fraction may appear in a very anodic position in Z(S) zone (in far left of this zone) for the following cases:
- blood sample with low Hb A level (< 10 %) and high Hb S level (for example, blood sample from transfused patient with sickle cell disease, or from patient with S beta-thalassaemia) for which the pattern is adjusted with Hb A and Hb A2 peaks, and,
 - blood sample without any Hb A and with high Hb S level for which the pattern is adjusted with Hb F and Hb A2 peaks.

This migration zone, which corresponds to an intermediary zone comprised within Z(S) zone, allows to automatically detect Hb S fractions with modified migration. It is called "Shifted Hb S area" and is indicated by a dash located on the left side of the name Z(S) in the upper section of the Z(S) migration zone. A variant which migrates in this position is identified as "Borderline variant" but not "Z(S) zone" like any other peak which migrates in the rest of the Z(S) zone.

Mix the sample with the Normal Hb A2 Control according to the procedure described in paragraph "SAMPLES FOR ANALYSIS", section "Particular cases", in order to confirm the position of the variant in the Z(S) migration zone. It is necessary to analyze the hematologic state and to perform complementary studies to check the presence of Hb S.

- When analyzing blood samples from patients with sickle cell disease before transfusion, a variation of Hb S fraction may be observed for the analyses of the same patient due to the inhomogeneity of this type of sample. It is therefore recommended to homogenize this type of blood sample before the analysis.

Interference and Limitations

- See SAMPLES FOR ANALYSIS.
- Analyze only blood samples contained in collection tubes indicated in the paragraph "EQUIPMENT AND ACCESSORIES REQUIRED" or tubes with equivalent dimensions approved for clinical assays. Call SEBIA technical service for further information on these devices.
- Do not analyze directly tubes containing less than 800 µL of blood sample.
- Avoid aged, improperly stored blood samples ; degradation products (or artefacts) may affect the electrophoretic pattern after 7 days storage.
- After 10 days storage, viscous aggregates composed in red blood cells may appear, they must be discarded before analysis.
- When analyzing blood samples with a decreased [red blood cells] / [plasma] ratio (from patients with anaemia), a hemoglobin Constant Spring variant may be suspected when a fraction is observed in Z(C) or Z(A2) migration zones. This fraction may be due to plasmatic proteins present in the sample (see § Sample preparation, Particular cases).
- When an abnormal hemoglobin is detected, use other means of identification (e.g., globin chain electrophoresis), or consult or send sample to a specialized laboratory.

IMPORTANT : It is also necessary to analyze the hematologic state, as complementary results.

- The migration of a hemoglobin variant close to Hb A involves an underestimation of Hb A fraction and that of the variant and consequently, an overestimation of Hb A2 fraction. In order to quantify Hb A2 with precision, it is necessary to delete the separate integration of both variants and Hb A, and to quantify these fractions together.
- Some homozygous "S" subjects receive a "Hydrea® (hydroxyurea) treatment that can induce synthesis of foetal hemoglobin. With CAPI 3 HEMOGLOBIN(E) procedure, the mobility of the induced hemoglobin F is not different from the physiological hemoglobin F.
- Due to the resolution and sensitivity limits of zone electrophoresis, it is possible that some hemoglobin variants may not be detected with this method.
- The CAPI 3 HEMOGLOBIN(E) procedure performed with the CAPILLARYS 3 instrument has not been evaluated in the neonate / newborn population (age range – birth to 28 days).
- In the case of patients with hyperleukocytosis, the migration speed of the sample may be accelerated causing a shift of the profile that may result in a non-recognition of the zones.

Hemoglobin variants observed with Hb A1c and / or HEMOGLOBIN(E) procedures :

Due to the different composition of Hb A1c and HEMOGLOBIN(E) buffers, the electrophoretic mobility of some hemoglobin variants may be different.

The common interfering factors with the CAPI 3 HEMOGLOBIN(E) procedure performed with the CAPILLARYS 3 instrument (triglycerides and bilirubin) were evaluated in studies based on the Clinical Laboratory Standards Institute (CLSI - USA) EP7-A2 guideline "Interference Testing in Clinical Chemistry".

The results are summarized below :

- No qualitative or quantitative interference with the CAPILLARYS 3 HEMOGLOBIN(E) procedure performed with the CAPILLARYS 3 instrument was detected if bilirubin concentration is equal to or less than 46.7 mg/dL, or 799 µmol/L.
- No qualitative or quantitative interference with the CAPILLARYS 3 HEMOGLOBIN(E) procedure performed with the CAPILLARYS 3 instrument was detected if triglycerides concentration is equal to or less than 2.3 g/dL, or 26.5 mmol/L.

Troubleshooting

Call SEBIA Technical Service of the supplier when the test fails to perform while the instruction for the preparation and storage of materials, and for the procedure were carefully followed.

Kit reagent Safety Data Sheets and information on i) cleaning and waste disposal, ii) labeling and safety rules applied by SEBIA, iii) packaging of biological samples for the transportation, and iv) instruments cleaning are available on the SEBIA's extranet website: www.sebia.com.

PERFORMANCE DATA

Precision

The precision of the CAPILLARYS 3 HEMOGLOBIN(E) procedure was evaluated in a study based on the Clinical and Laboratory Standards Institute (CLSI - USA) EP5-A2 guideline "Evaluation of Precision Performance of Quantitative Measurement Methods; Approved Guideline – Second Edition".

The means and coefficients of variation (CV %) were calculated for percentage (%) of hemoglobin fractions for each sample, using statistical tools recommended by CLSI.

Reproducibility within the same capillary from the same instrument

Twelve (12) different blood samples were run using the CAPILLARYS 3 HEMOGLOBIN(E) procedure on the CAPILLARYS 3 instrument.

In this study, each blood sample was analyzed on the same capillary from the same instrument and with 3 lots of CAPILLARYS 3 HEMOGLOBIN(E) kit, including 6 runs over 3 working days (at 2 different times of the day). Within each run, samples were analyzed in duplicate.

CV ranges were obtained for each fraction by conducting this study on all capillaries from 3 different instruments.

The following table summarizes the repeatability and the total reproducibility CV (%) ranges for percentages (%) of each hemoglobin fraction from all samples.

Fraction	Ranges of % tested		Repeatability						Total reproducibility					
			Instrument No. 1		Instrument No. 2		Instrument No. 3		Instrument No. 1		Instrument No. 2		Instrument No. 3	
	Min value	Max value	CV min (%)	CV max (%)	CV min (%)	CV max (%)	CV min (%)	CV max (%)	CV min (%)	CV max (%)	CV min (%)	CV max (%)	CV min (%)	CV max (%)
Hb A	28.4	98.3	0.0	0.7	0.0	0.7	0.0	1.0	0.0	0.8	0.0	0.8	0.0	1.4
Hb A2	1.7	7.3	0.0	4.0	0.0	3.4	0.0	3.5	0.0	4.4	0.0	3.9	0.0	5.1
Hb F	3.6	69.0	0.1	1.8	0.0	1.9	0.0	1.6	0.2	2.4	0.0	2.3	0.1	2.3
Hb S	17.5	33.9	0.1	0.7	0.1	0.6	0.1	1.1	0.3	1.2	0.3	1.0	0.3	1.4
Hb C	6.8	33.5	0.0	1.5	0.0	1.0	0.4	1.0	0.0	1.5	0.0	1.3	0.7	2.4
Hb D	40.2		0.1	0.3	0.1	0.4	0.1	0.5	0.2	0.4	0.2	0.5	0.4	0.8
Hb E	21.8		0.2	0.5	0.2	0.7	0.1	0.9	0.4	1.0	0.4	0.9	0.4	3.1

Reproducibility between capillaries from the same instrument

Twelve (12) different blood samples were run using the CAPILLARYS 3 HEMOGLOBIN(E) procedure on the CAPILLARYS 3 instrument.

In this study, each blood sample was analyzed on all capillaries from the same instrument and with 1 lot of CAPILLARYS 3 HEMOGLOBIN(E) kit, including 12 runs over 6 working days (at 2 different times of the day). Within each run, samples were analyzed in quadruplicate.

CV ranges were obtained for each fraction by conducting this study with 3 lots of kit on 3 different instruments.

The following table summarizes the repeatability and the total reproducibility CV (%) ranges for percentages (%) of each hemoglobin fraction from all samples.

Fraction	Ranges of % tested		Repeatability						Total reproducibility					
			Instrument No. 1		Instrument No. 2		Instrument No. 3		Instrument No. 1		Instrument No. 2		Instrument No. 3	
	Min value	Max value	CV min (%)	CV max (%)	CV min (%)	CV max (%)	CV min (%)	CV max (%)	CV min (%)	CV max (%)	CV min (%)	CV max (%)	CV min (%)	CV max (%)
Hb A	28.4	98.3	0.0	0.5	0.0	0.5	0.0	0.7	0.0	0.7	0.0	0.7	0.0	1.2
Hb A2	1.7	7.3	0.7	3.0	0.7	3.3	0.5	3.1	0.9	4.5	0.8	4.3	0.6	5.1
Hb F	3.6	69.0	0.2	2.2	0.2	2.1	0.3	1.6	0.3	2.3	0.3	3.1	0.3	1.8
Hb S	17.5	33.9	0.4	0.8	0.3	0.5	0.4	0.8	0.5	0.8	0.5	0.7	0.5	1.0
Hb C	6.8	33.5	0.5	0.9	0.6	0.9	0.6	1.1	0.8	1.1	0.8	1.1	0.8	2.2
Hb D	40.2		0.2	0.3	0.2	0.3	0.3	0.5	0.3	0.4	0.3	0.4	0.4	0.6
Hb E	21.8		0.4	0.5	0.5	0.7	0.6	1.6	0.5	0.8	0.6	0.8	0.7	1.8

Reproducibility between lots and between instruments

Twelve (12) different blood samples were run using the CAPI 3 HEMOGLOBIN(E) procedure on the CAPILLARYS 3 instrument.

In this study, each blood sample was analyzed at 2 different times of the day on all capillaries from 3 different instruments and with 3 lots of CAPI 3 HEMOGLOBIN(E) kit. Within each run, samples were analyzed in quadruplicate.

The analysis of obtained results allows to demonstrate the reproducibility :

- between lots : from data obtained with 3 lots of CAPI 3 HEMOGLOBIN(E) kit on the same instrument, including 36 runs over 18 working days. CV ranges were obtained for each fraction by conducting this study on 3 different instruments.
- between instruments : from data obtained with 3 instruments and 1 lot of CAPI 3 HEMOGLOBIN(E) kit, including 36 runs over 18 working days. CV ranges were obtained for each fraction by conducting this study on 3 different lots.
- between lots and between instruments : from combined data obtained with the 3 instruments and the 3 lots of CAPI 3 HEMOGLOBIN(E) kit, including 108 runs over 54 working days.

The following table summarizes the repeatability and the total reproducibility CV (%) ranges for percentages (%) of each hemoglobin fraction from all samples.

Fraction	Ranges of % tested		Reproducibility between lots				Reproducibility between instruments				Reproducibility between lots and between instruments			
			Repeatability		Total reproducibility		Repeatability		Total reproducibility		Repeatability		Total reproducibility	
	Min value	Max value	CV min (%)	CV max (%)	CV min (%)	CV max (%)	CV min (%)	CV max (%)	CV min (%)	CV max (%)	CV min (%)	CV max (%)	CV min (%)	CV max (%)
Hb A	28.4	98.3	0.0	0.6	0.0	0.8	0.0	0.6	0.0	1.0	0.0	0.5	0.0	0.9
Hb A2	1.7	7.3	0.7	3.0	0.9	4.9	0.7	3.1	1.0	5.2	0.8	2.9	1.1	5.0
Hb F	3.6	69.0	0.2	1.9	0.3	2.5	0.2	1.8	0.3	2.4	0.2	1.7	0.4	2.3
Hb S	17.5	33.9	0.4	0.6	0.5	0.8	0.4	0.7	0.6	0.9	0.5	0.5	0.8	0.8
Hb C	6.8	33.5	0.7	0.9	0.9	1.5	0.7	0.9	0.9	1.4	0.8	0.8	1.1	1.2
Hb D	40.2		0.3	0.4	0.4	0.5	0.3	0.3	0.3	0.4	0.3		0.4	
Hb E	21.8		0.5	1.1	0.7	1.3	0.5	1.0	0.8	1.2	0.8		1.0	

Linearity

Mixture of 2 different blood samples

This linearity study of the CAPI 3 HEMOGLOBIN(E) procedure was evaluated in a study based on the Clinical and Laboratory Standards Institute (CLSI - USA) EP6-A guideline "Evaluation of the Linearity of Quantitative Measurement Procedures: A Statistical Approach; Approved Guideline".

The results for percentage (%) of hemoglobin fractions were analyzed using statistical tools recommended by CLSI.

Hb A & Hb S fractions

2 characteristic blood samples, including a blood sample with normal Hb A2 level (containing 14.1 g/dL total hemoglobin with 0.0 % Hb S and 97.3 % Hb A) and a blood sample with Hb S (containing 7.9 g/dL total hemoglobin with 89.7 % Hb S and 0.0 % Hb A) were mixed within different proportions and the mixtures were electrophoresed with CAPI 3 HEMOGLOBIN(E) procedure performed with CAPILLARYS 3 instrument. For each mixture, samples were analyzed in triplicate.

The tests were determined to be linear within the entire range studied for Hb A fraction until a maximum concentration of about 13.7 g/dL (between 0.9 and 97.3 % Hb A) and for Hb S fraction until a maximum concentration of about 7.1 g/dL (between 0.8 and 89.7 % Hb S).

Hb A2 fraction

2 characteristic blood samples, including a Hb A2 depleted blood sample (containing 13.6 g/dL total hemoglobin with 0.0 % Hb A2) and a Hb A2 enriched blood sample (containing 14.4 g/dL total hemoglobin with 9.1 % Hb A2) were mixed within different proportions and the mixtures were electrophoresed with CAPI 3 HEMOGLOBIN(E) procedure performed with CAPILLARYS 3 instrument. For each mixture, samples were analyzed in triplicate.

The tests were determined to be linear within the entire range studied for Hb A2 fraction until a maximum concentration of about 1.3 g/dL (between 0.2 and 9.1 % Hb A2).

Hb F fraction

2 characteristic blood samples, including a blood sample with normal Hb A2 level (containing 13.6 g/dL total hemoglobin with 0.0 % Hb F) and a blood sample with increased Hb F level (containing 13.7 g/dL total hemoglobin with 83.1 % Hb F) were mixed within different proportions and the mixtures were electrophoresed with CAPI 3 HEMOGLOBIN(E) procedure performed with CAPILLARYS 3 instrument. For each mixture, samples were analyzed in triplicate.

The tests were determined to be linear within the entire range studied for Hb F fraction until a maximum concentration of about 11.4 g/dL (between 0.5 and 83.1 % Hb F).

Hb C fraction

2 characteristic blood samples, including a blood sample with normal Hb A2 level (containing 12.9 g/dL total hemoglobin with 0.0 % Hb C) and a blood sample with Hb C (containing 9.3 g/dL total hemoglobin with 82.0 % Hb C) were mixed within different proportions and the mixtures were electrophoresed with CAPI 3 HEMOGLOBIN(E) procedure performed with CAPILLARYS 3 instrument. For each mixture, samples were analyzed in triplicate.

The tests were determined to be linear within the entire range studied for Hb C fraction until a maximum concentration of about 7.6 g/dL (between 0.3 and 82.0 % Hb C).

Hb D fraction

2 characteristic blood samples, including a blood sample with normal Hb A2 level (containing 16.4 g/dL total hemoglobin with 0.0 % Hb D) and a blood sample with Hb D (containing 12.7 g/dL total hemoglobin with 43.5 % Hb D) were mixed within different proportions and the mixtures were electrophoresed with CAPI 3 HEMOGLOBIN(E) procedure performed with CAPILLARYS 3 instrument. For each mixture, samples were analyzed in triplicate.

The tests were determined to be linear within the entire ranges studied for Hb D fraction until a maximum concentration of about 5.5 g/dL (between 0.7 and 43.5 % Hb D).

Hb E fraction

2 characteristic blood samples, including a blood sample with normal Hb A2 level (containing 12.5 g/dL total hemoglobin with 0.0 % Hb E) and a blood sample with Hb E (containing 8.8 g/dL total hemoglobin with 86.9 % Hb E) were mixed within different proportions and the mixtures were electrophoresed with CAPI 3 HEMOGLOBIN(E) procedure performed with CAPILLARYS 3 instrument. For each mixture, samples were analyzed in triplicate.

The tests were determined to be linear within the entire ranges studied for Hb E fraction until a maximum concentration of about 7.6 g/dL (between 0.2 and 86.9 % Hb E).

Dilution in hemolysing solution**Hb A & Hb F fractions**

A blood sample with increased Hb F level (containing 10.5 g/dL total hemoglobin with 18.7 % Hb A and 81.3 % Hb F) was concentrated and serially diluted in hemolysing solution and the mixtures were electrophoresed with the CAPI 3 HEMOGLOBIN(E) procedure performed with CAPILLARYS 3 instrument.

The tests were determined to be linear within the entire ranges studied from 1.1 to 21.0 g/dL total hemoglobin and Hb A and Hb F fraction percentages were not affected by the hemoglobin concentration of the samples.

Hb A2 fraction

A blood sample with normal Hb A2 level (containing 9.2 g/dL total hemoglobin with 2.6 % Hb A2) was concentrated and serially diluted in hemolysing solution and the mixtures were electrophoresed with the CAPI 3 HEMOGLOBIN(E) procedure performed with CAPILLARYS 3 instrument.

The tests were determined to be linear within the entire ranges studied from 1.8 to 21.9 g/dL total hemoglobin and Hb A2 fraction percentages were not affected by the hemoglobin concentration of the samples.

Hb S fraction

A blood sample with Hb S (containing 12.5 g/dL total hemoglobin with 40.7 % Hb S) was concentrated and serially diluted in hemolysing solution and the mixtures were electrophoresed with the CAPI 3 HEMOGLOBIN(E) procedure performed with CAPILLARYS 3 instrument.

The tests were determined to be linear within the entire range studied from 2.5 to 20.9 g/dL total hemoglobin and Hb S fraction percentages were not affected by the hemoglobin concentration of the samples.

Hb C fraction

A blood sample with Hb C (containing 11.9 g/dL total hemoglobin with 31.5 % Hb C) was concentrated and serially diluted in hemolysing solution and the mixtures were electrophoresed with the CAPI 3 HEMOGLOBIN(E) procedure performed with CAPILLARYS 3 instrument.

The tests were determined to be linear within the entire ranges studied from 2.4 to 19.9 g/dL total hemoglobin and Hb C fraction percentages were not affected by the hemoglobin concentration of the samples.

Hb D fraction

A blood sample with Hb D (containing 9.7 g/dL total hemoglobin with 43.2 % Hb D) was concentrated and serially diluted in hemolysing solution and the mixtures were electrophoresed with the CAPI 3 HEMOGLOBIN(E) procedure performed with CAPILLARYS 3 instrument.

The tests were determined to be linear within the entire ranges studied from 1.9 to 19.4 g/dL total hemoglobin and Hb D fraction percentages were not affected by the hemoglobin concentration of the samples.

Hb E fraction

A blood sample with Hb E (containing 8.9 g/dL total hemoglobin with 24.5 % Hb E) was concentrated and serially diluted in hemolysing solution and the mixtures were electrophoresed with the CAPI 3 HEMOGLOBIN(E) procedure performed with CAPILLARYS 3 instrument.

The tests were determined to be linear within the entire ranges studied from 1.8 to 21.2 g/dL total hemoglobin and Hb E fraction percentages were not affected by the hemoglobin concentration of the samples.

Accuracy – Internal correlation

The internal concordance study of the CAPI 3 HEMOGLOBIN(E) procedure was evaluated in a study based on the Clinical and Laboratory Standards Institute (CLSI - USA) EP09-A2-IR guideline "Method Comparison and Bias Estimation Using Patient Samples; Approved Guideline – Second Edition (Interim Revision)".

The results for percentages (%) of hemoglobin fractions were analyzed using statistical tools recommended by CLSI.

NOTE : The results presented below have been obtained from 1 internal accuracy study. The analyzed blood samples were provided by 10 laboratories in France, Belgium, Thailand and New Zealand. All the samples were exactly treated the same way with both techniques and followed the same guidelines in regards to sample integrity.

The levels of hemoglobin fractions were measured in 153 blood samples, including 64 samples with hemoglobin variants, both by electrophoretic separations obtained with the CAPI 3 HEMOGLOBIN(E) procedure performed with the CAPILLARYS 3 instrument and a commercially available capillary electrophoresis technique for hemoglobin analysis (reference).

The measured values of hemoglobin fractions from both procedures were analyzed by a linear regression statistical procedure. Sensibility and specificity of the CAPI 3 HEMOGLOBIN(E) procedure compared to the reference procedure have been calculated using the recommended method (Wendling, 1986). The results of linear regression analysis are tabulated below ($y = \text{CAPI 3 HEMOGLOBIN(E) with CAPILLARYS 3 instrument}$) :

Normal hemoglobins

Fraction	Number of samples	Correlation coefficient	y-intercept	Slope	Range of Hb % values CAPI 3 HEMOGLOBIN(E)	Sensitivity (%)	Specificity (%)
Hb A	150	1.000	-0.993	1.010	16.9 - 98.7	100.0	100.0
Hb A2	148	0.998	0.005	0.986	0.5 - 9.2	100.0	100.0
Hb F	22	1.000	-0.008	1.009	0.8 - 83.1	100.0	100.0

Hemoglobin variants

Fraction	Number of samples	Correlation coefficient	y-intercept	Slope	Range of Hb % values CAPI 3 HEMOGLOBIN(E)
Hb S	13	1.000	-0.025	1.010	1.8 - 89.7
Hb C	13	1.000	0.099	1.008	2.0 - 89.5
Hb D	9	1.000	-0.068	1.015	3.3 - 43.7
Hb E	13	1.000	0.183	1.001	5.0 - 86.9

This study demonstrated a perfect correlation between the 2 analysis procedures for the Hb A, Hb A2, Hb F, Hb S, Hb C, Hb D and Hb E quantitative determination.

For the detection of hemoglobin variants, the obtained results demonstrate a perfect correlation between the 2 analysis procedures, with a 100.0 % sensibility and a 100.0 % specificity of CAPI 3 HEMOGLOBIN(E) procedure compared to the reference procedure.

All abnormal hemoglobins or abnormal levels of normal hemoglobins detected with the CAPI 3 HEMOGLOBIN(E) procedure performed with the CAPILLARYS 3 instrument were in agreement with the reference procedure. There was no case observed of false positive, i.e., detection of an abnormal band or abnormal level of a normal band where no such abnormality existed.

Limit of blank (LOB) – Limit of detection (LOD)

The determination of the limit of blank (LOB) and the limit of detection (LOD) of the CAPI 3 HEMOGLOBIN(E) procedure was evaluated in a study based on the Clinical and Laboratory Standards Institute (CLSI - USA) EP17-A guideline "Protocols for Determination of Limits of Detection and Limits of Quantitation; Approved Guideline".

For each hemoglobin fraction, the limit of blank (LOB) is determined using 5 different blood samples and the limit of detection (LOD) is determined using 5 different blood samples.

The results are tabulated below :

Fraction	LOB (%)	LOD (%)
Hb A	0.1	0.9
Hb A2	0.1	0.2
Hb F	0.1	0.5
Hb S	0.2	0.8
Hb C	0.1	0.3
Hb D	0.1	0.7
Hb E	0.1	0.2

BIBLIOGRAPHIE / BIBLIOGRAPHY

BIBLIOGRAFIE - BIBLIOGRAFIA - BIBLIOGRAFÍA - BIBLIOGRAFI - ΒΙΒΛΙΟΓΡΑΦΙΑ - BIBLIOGRAFIJU - BIBLIOGRAFIJA - КАРНАҚА - БИБЛІОГРАФІЯ - 参考书目 - БИБЛІОГРАФІЮ - 参考文献 - İZMANTOTÂ LİTERATÜRA - BIBLIOGRAFIU - KIRJANDUS - ДАНН МҮС ТАІ ЛІЕҮ ТҘМ ҚҘО

1. J. Bardakdjian-Michau, J.-L. Dhondt, R. Ducrocq, F. Galactéros, A. Guyard, F.-X. Huchet, A. Lahary, D. Lena-Russo, P. Maboudou, M.-L. North, C. Prehu, A.-M. Soummer, M. Verschelde, H. Wajzman (2003) Bonnes pratiques de l'étude de l'hémoglobine. *Ann. Biol. Clin.*, 61, 401-409.
2. V.F. Fairbanks, ed. (1980) Hemoglobinopathies and thalassemia: Laboratory methods and case studies. Brian C. Decker, New York.
3. F. Galacteros (1986) Thalassémie, drépanocytose et autres hémoglobinopathies. *Techniques et Biologie*, 3, 174-178.
4. JM Hempe, JN Granger and RD Craver (1997) Capillary isoelectric focusing of hemoglobin variants. *Electrophoresis*, 18, 1785-1795.
5. T.H.J. Huisman and J.H.P. Jonxis (1977) The hemoglobinopathies: techniques of identification. Marcel Dekker, New York.
6. Jellum E *et al.* Diagnostic applications of chromatography and capillary electrophoresis. *J. Chromatogr. B*, 689, 155-164 (1997).
7. Joutovsky A, Hadzi-Nesic J and Nardi MA (2004) HPLC retention time as a diagnostic tool for hemoglobin variants and hemoglobinopathies : a study of 60 000 samples in a clinical diagnostic laboratories. *Clin. Chem.*, 50, 10, 1736-1747.
8. J.S. Krauss, P.A. Drew, M.H. Jonah, M. Trinh, S. Shell, L. Black and C.R. Baisden (1986) Densitometry and microchromatography compared for determination of the hemoglobin C and A2 proportions in hemoglobin C and hemoglobin SC disease and in hemoglobin C trait. *Clin. Chem.* 32, 5, 860-863.
9. Landers JP. Clinical Capillary Electrophoresis. *Clin. Chem.*, 41, 495-509 (1995).
10. C. Livingstone (1986) The hemoglobinopathies. Edit. London.-
11. M. Maier-Redelsberger, R. Girot (1989) Diagnostic biologique des maladies de l'hémoglobine. *Feuilles de biologie*, 170.
12. Oda RP *et al.* Capillary electrophoresis as a clinical tool for the analysis of protein in serum and other body fluids. *Electrophoresis*, 18, 1715-1723 (1997).
13. R.G. Schneider (1978) Methods for detection of hemoglobin variants and hemoglobinopathies in the routine clinical laboratory. *CRC Crit. Rev. Clin. Lab. Sci.* 9, 243-271.
14. L. Vovan, D. Lara-Russo, A. Orsini (1985) Diagnostic biologique des hémoglobinoses. *Ann. Pédiat.* 32, 9, 780-789.
15. <http://globin.cse.psu.edu/hbvar/menu.html> : Hbvar : A Database of Human Hemoglobin Variants and Thalassemias.
16. <http://www.isns-neoscreening.org/agenda.htm>.
17. F Boemer, O Ketelslegers, JM Miron, V Bours, R Schoops (2008) Newborn screening for sickle cell disease using tandem mass spectrometry. *Clin. Chem.*, 54, 12, 2036-2041.
18. Lubin BH, Witkowska HE, Kleman K (1991) Laboratory diagnosis of hemoglobinopathies. *Clin. Biochem.*, 24, 363-374.
19. B Gulbis, B Fontaine, F Vertongen, F Cotton (2003) The place of capillary electrophoresis techniques in screening for haemoglobinopathies. *Ann. Clin. Biochem.*, 40, 659-662.
20. Aguilar-Martinez P *et al* (2010) Arbres décisionnels pour le diagnostic et la caractérisation moléculaire des hémoglobinopathies. *Ann. Biol. Clin.*, 68 (4) : 455-464.
21. Wendling A. Procédures de diagnostic ou de dépistage : Justification et validité d'un test de diagnostic ou de dépistage-sensibilité-spécificité. *Impact-Internat*, 1986 ; Sept : 93-97.
22. L. Guis, A. Chaumier, V. Le Gall, S. Havrez (Février 2013) Intégration du Capillarys 2 Flex Piercing (Sebia) dans un laboratoire de biologie médicale spécialisée. *Revue Francophone des Laboratoires*, 449, 47 – 56.
23. I. Agouti, F. Merono, N. Bonello-Palot, C. Badens (2013) Analytical evaluation of the Capillarys 2 Flex piercing for routine haemoglobinopathies diagnosis. *Int. Jnl. Lab. Hem.* 35, 217 – 221.
24. S. Altinier, M. Varagnolo, M. Zaninotto, M. Plebani. (2012) Identification and quantification of hemoglobins in whole blood: the analytical and organizational aspects of Capillarys 2 Flex Piercing compared with agarose electrophoresis and HPLC methods. *Clin. Chem. Lab. Med.* DOI 10.1515/ccim-2012-0061.
25. M. Agastiniotis, J.L. Vives Corrons, E.S. Soteriades, A. Eleftheriou (2013) The Impact of Migrations on the Health Services for Rare Diseases in Europe: The Example of Haemoglobin Disorders. *The Scientific World Journal* Volume 2013, Article ID 727905, 10 pages.
26. Nicole Borbely, Lorraine Phelan, Richard Szydlo, *et al.* (2012) Capillary zone electrophoresis for haemoglobinopathy diagnosis. *J. Clin. Pathol.*, doi: 10.1136/jclinpath-2012-200946.
27. F. Cotton *et al.* (2009) Evaluation of an automated capillary electrophoresis system in the screening for hemoglobinopathies. *Clin. Lab.*, 55 : 217 – 221.
28. D. Greene, A.L. Pyle, J.S. Chang, C. Hoke, T. Lorey (2012) Comparison of Sebia Capillarys Flex capillary electrophoresis with the BioRad Variant II high pressure liquid chromatography in the evaluation of hemoglobinopathies. *Clin. Chim. Acta* 413, 1232 – 1238.
29. T. Higgins, M. Maek, A. Khajuria (2009) Comparison of two methods for the quantification and identification of hemoglobin variants. *Clin. Biochem.*, 42, 701 – 705.
30. D.F. Keren *et al* (2008) Comparison of Sebia Capillarys capillary electrophoresis with the Primus High-Pressure Liquid Chromatography in the evaluation of hemoglobinopathies. *Am. J. Clin. Pathol.*, 130 : 824 – 831.
31. D.F. Keren *et al* (2012) Expression of hemoglobin variant migration by capillary electrophoresis relative to Hemoglobin A2 improves precision. *Am. J. Clin. Pathol.*, 137 : 660 – 664.
32. Can Liao, Jian-Ying Zhou, Xing-Mei Xie, Jian Li, Ru Li, and Dong-Zhi Li (2010). Detection of Hb constant spring by a capillary electrophoresis method. *Hemoglobin*, 34 (2) : 175 – 178.
33. D.M. Mais *et al* (2009) The range of hemoglobin A2 in hemoglobin E heterozygotes as determined by capillary electrophoresis. *Am. J. Clin. Pathol.*, 132 : 34 – 38.
34. T. Munksgaard *et al* (2010) Quantitative analysis of Hb Bart's in cord blood by capillary electrophoresis system. *Ann. Hematol.* DOI 10.1007/s00277-010-1137-4.
35. R. Paleari, B. Gulbis, F. Cotton, A. Mosca (2012) Interlaboratory comparison of current high-performance methods for HbA2. *Int. Jnl. Lab. Hem.* 34, 362 – 368.
36. S. Sangkitporn *et al* (2011) Multicenter validation of fully automated capillary electrophoresis method for diagnosis of thalassemias and hemoglobinopathies in Thailand. *Southeast Asian J. Trop. Med. Public Health* 42 (5), 1224 – 1232.
37. F. Wolff, F. Cotton, B. Gulbis (2012) Screening for haemoglobinopathies on cord blood : laboratory and clinical experience. *J. Med. Screen.*, 19, 3, 116 - 122.

BIBLIOGRAPHIE / BIBLIOGRAPHY

BIBLIOGRAFIE - BIBLIOGRAFIA - BIBLIOGRAFÍA - BIBLIOGRAFI - ΒΙΒΛΙΟΓΡΑΦΙΑ - BIBLIOGRAFIJU - BIBLIOGRAFIJA - КАРНАҚА - БИБЛИОГРАФИЯ - 参考书目 - БИБЛИОГРАФИЮ - 参考文献 - IZMANTOTĀ LĪTERĀTŪRA - BIBLIOGRAFIU - KIRJANDUS - DANH MỤC TÀI LIỆU THAM KHẢO

38. A.D. Stephens *et al* on behalf of the International Council for the Standardisation of Haematology (ICSH) (2012). ICSH recommendations for the measurement of Haemoglobin F. *Int. Jnl. Lab. Hem.*, 34, 14 – 20.
39. A. Mosca, R. Paleari, D. Leone, G. Ivaldi (2009) The relevance of hemoglobin F measurement in the diagnosis of thalassemias and related hemoglobinopathies. *Clinical Biochemistry*, 42, 1797–1801.
40. J. D. Hoyer, C. S. Penz, V. F. Fairbanks, C. A. Hanson and J.A. Katzmann (2002) Diagnostic Usefulness in the Distinction of Hereditary Persistence of Fetal Hemoglobin (HPFH) and Hemoglobin S–HPFH From Other Conditions With Elevated Levels of Hemoglobin F. *Am. J. Clin. Pathol.*, 117 : 857 – 863.
41. P. Van Delft *et al* (2009) Evaluating five dedicated automatic devices for haemoglobinopathy diagnostics in multi-ethnic populations. *Int. J. Lab. Hematol.*, 31 (5) : 484 – 95.
42. Prevention and diagnosis of haemoglobinopathies. A short guide for health professionals and laboratory scientists. Thalassemia International Federation (TIF) publication No. 21.
43. A.J. Marengo-Rowe (2007) The thalassemias and related disorders. *Baylor University Medical Center Proceedings*, 20, 27 - 31.
44. Sae-ung *et al* (2012). Phenotypic expression of hemoglobins A2, E and F in various hemoglobin E related disorders. *Blood Cells, Molecules, and Diseases*, 48, 11–16.
45. A. Cao and R. Galanello (2010) Beta-thalassemia. *Genetics in Medicine*, 12 (2) : 61 – 76.
46. R.Z. Azma *et al* (2012) Co-inheritance of compound heterozygous Hb Constant Spring and a single – α 3.7 gene deletion with heterozygous $\delta\beta$ thalassaemia: A diagnostic challenge. *Malaysian J. Pathol.* ; 34(1) : 57 – 62.
47. C.L. Harteveld and D.R. Higgs (2010) α -thalassaemia. *Orphanet Journal of Rare Diseases*, 5, 13.

TABLEAU / TABLE

**CAP1 3 HEMOGLOBIN(E) : VARIANTS POTENTIELS PRÉSENTS DANS CHAQUE ZONE
POTENTIAL VARIANTS LOCATED IN EACH ZONE**

Zone	Hémoglobines / Hemoglobins (Hb)
Z1	<p>Hb Santa Ana (pic mineur) #, Hb Mizuho (pic mineur) #, Hb delta A'2, Hb A2-Canebière, Hb A2-Lampang, Hb S-Oman, Hb A2-Turkish, Hb T-Cambodia, Hb Poissy (pic mineur) #, variant de Hb A2 "Chad" #, variant de Hb A2 "Savaria" #, variant de Hb A2 "Arya" #, variant de Hb A2 "Hasharon" #, variant de Hb A2 "Fort de France" #, variant de Hb A2 "Ottawa" #, variant de Hb A2 "Shimonoseki" #, variant de Hb A2 "Russ" (alpha 2) #, variant de Hb A2 "Russ" (alpha 1) #, variant de Hb A2 "Matsue-OkI" #, variant de Hb A2 "Reims" #, variant de Hb A2 "Mizushi" #, variant de Hb A2 "Stanleyville-II" #, variant de Hb A2 "O-Indonesia" #, variant de Hb A2 "San Antonio" #, variant de Hb A2 "G-Audhali" #, variant de Hb A2 "Handsworth" #, variant de Hb A2 "G-Philadelphia" #, variant de Hb A2 "Q-India" #, variant de Hb A2 "Memphis" #, variant de Hb A2 "Q-Iran" #, variant de Hb A2 "G-Waimanalo" #, variant de Hb A2 "Watts" #, variant de Hb A2 "Spanish Town" #, variant de Hb A2 "Montgomery" #, variant de Hb A2 "G-Norfolk" #, variant de Hb A2 "Inkster" #, variant de Hb A2 "Ube-4" #, variant de Hb A2 "G-Pest" #, variant de Hb A2 "Winnipeg" #, variant de Hb A2 "Queens" #, variant de Hb A2 "Etobicoke" #, variant de Hb A2 "Chapel Hill" #, variant de Hb A2 "Park Ridge" #, variant de Hb A2 "Q-Thailand" #, variant de Hb A2 "Delfzicht" # !!</p> <p><i>Hb Santa Ana (minor peak) #, Hb Mizuho (minor peak) #, Hb delta A'2, Hb A2-Canebière, Hb A2-Lampang, Hb S-Oman, Hb A2-Turkish, Hb T-Cambodia, Hb Poissy (minor peak) #, "Chad" Hb A2 variant #, "Savaria" Hb A2 variant #, "Arya" Hb A2 variant #, "Hasharon" Hb A2 variant #, "Fort de France" Hb A2 variant #, "Ottawa" Hb A2 variant #, "Shimonoseki" Hb A2 variant #, "Russ" Hb A2 variant (alpha 2) #, "Russ" Hb A2 variant (alpha 1) #, "Matsue-OkI" Hb A2 variant #, "Reims" Hb A2 variant #, "Mizushi" Hb A2 variant #, "Stanleyville-II" Hb A2 variant #, "O-Indonesia" Hb A2 variant #, "San Antonio" Hb A2 variant #, "G-Audhali" Hb A2 variant #, "Handsworth" Hb A2 variant #, "G-Philadelphia" Hb A2 variant #, "Q-India" Hb A2 variant #, "Memphis" Hb A2 variant #, "Watts" Hb A2 variant #, "Spanish Town" Hb A2 variant #, "Montgomery" Hb A2 variant #, "G-Norfolk" Hb A2 variant #, "Inkster" Hb A2 variant #, "Ube-4" Hb A2 variant #, "G-Pest" Hb A2 variant #, "Winnipeg" Hb A2 variant #, "Queens" Hb A2 variant #, "Etobicoke" Hb A2 variant #, "Chapel Hill" Hb A2 variant #, "Park Ridge" Hb A2 variant #, "Q-Thailand" Hb A2 variant #, "Delfzicht" Hb A2 variant # !!</i></p>
Z(C)	<p>Hb C-Ziguinchor !!, Hb F-Hull, Hb F-Texas-I, Hb Constant Spring, Hb Paksé, Hb C, Hb C-Harlem (C-Georgetown), variant de Hb A2 "Les Lilas" #, variant de Hb A2 "Boumerdes" #, variant de Hb A2 "Tarrant" #, variant de Hb A2 "Dunn" #, variant de Hb A2 "Bassett" #, variant de Hb A2 "Sassari" #, variant de Hb A2 "St. Luke's" #, variant de Hb A2 "Verdun" #, variant de Hb A2 "Manitoba-I" #, variant de Hb A2 "Setif" #, variant de Hb A2 "Sunshine Seth" #, variant de Hb A2 "Titusville" #, variant de Hb A2 "Swan River" #, variant de Hb A2 "Manitoba-II" #, variant de Hb A2 "Val de Marne" #</p> <p><i>Hb C-Ziguinchor !!, Hb F-Hull, Hb F-Texas-I, Hb Constant Spring, Hb Paksé, Hb C, Hb C-Harlem (C-Georgetown), "Les Lilas" Hb A2 variant #, "Boumerdes" Hb A2 variant #, "Tarrant" Hb A2 variant #, "Dunn" Hb A2 variant #, "Bassett" Hb A2 variant #, "Sassari" Hb A2 variant #, "St. Luke's" Hb A2 variant #, "Verdun" Hb A2 variant #, "Manitoba-I" Hb A2 variant #, "Setif" Hb A2 variant #, "Sunshine Seth" Hb A2 variant #, "Titusville" Hb A2 variant #, "Swan River" Hb A2 variant #, "Manitoba-II" Hb A2 variant #, "Val de Marne" Hb A2 variant #</i></p>
Z(A2)	<p>Hb A2, Hb Chad (E-Keelung) #, Hb A2-Madrid * #, Hb A2-Saint Denis * #, Hb A2-Saint-Etienne * #, Hb Hong Kong (cas anti-Lepore), Hb O-Tibesti, Hb Gun Hill, Hb O-Arab, Hb E-Saskatoon, Hb Shuangfeng, variant de Hb A2 "Charolles" #, variant de Hb A2 "Roubaix" #, variant de Hb A2 "El Escorial" * #, variant de Hb A2 "Dallas" * #, variant de Hb A2 "Barika" * #, variant de Hb A2 "Melusine" * #, variant de Hb A2 "Jura" #, variant de Hb A2 "Nouakchott" #, variant de Hb A2 "Pohnpei" #</p> <p><i>Hb A2, Hb Chad (E-Keelung) #, Hb A2-Madrid * #, Hb A2-Saint Denis * #, Hb A2-Saint-Etienne * #, Hb Hong Kong (anti-Lepore case), Hb O-Tibesti, Hb Gun Hill, Hb O-Arab, Hb E-Saskatoon, Hb Shuangfeng, "Charolles" Hb A2 variant #, "Roubaix" Hb A2 variant #, "El Escorial" Hb A2 variant * #, "Dallas" Hb A2 variant * #, "Barika" Hb A2 variant * #, "Melusine" Hb A2 variant * #, "Jura" Hb A2 variant #, "Nouakchott" Hb A2 variant #, "Pohnpei" Hb A2 variant #</i></p>
Z(E)	<p>Hb Hornchurch, Hb Seal Rock, Hb Köln (Ube-1) #, Hb Buenos Aires (pic mineur) #, Hb E, Hb Cleveland, Hb M-Saskatoon (pic mineur) #, Hb G-Siriraj, Hb A2-Babinga, Hb F-Moyen Orient, Hb O-Padova, Hb Agenogi, Hb Sabine #, Hb Santa Ana #, Hb Savaria # !!, Hb Djelfa (pic 3) # !!, variant de Hb A2 "M-Iwate" #, variant de Hb A2 "Saint Claude" #, variant de Hb A2 "Jackson" (alpha 2) #, Hb C dégradée</p> <p><i>Hb Hornchurch, Hb Seal Rock, Hb Köln (Ube-1) #, Hb Buenos Aires (minor peak) #, Hb E, Hb Cleveland, Hb M-Saskatoon (minor peak) #, Hb G-Siriraj, Hb A2-Babinga, Hb F-Moyen Orient, Hb O-Padova, Hb Agenogi, Hb Sabine #, Hb Santa Ana #, Hb Savaria # !!, Hb Djelfa (peak 3) # !!, "M-Iwate" Hb A2 variant #, "Saint Claude" Hb A2 variant #, "Jackson" Hb A2 variant (alpha 2) #, denatured Hb C</i></p>
Z(S)	<p>Hb Arya # !!, Hb Kenya (HPFH-7), Hb Hasharon (Sinai) #, Hb Dhofar (Yukuhashi), Hb Shimonoseki (Hikoshima) #, Hb O-Indonesia (Buginese-X) #, Hb Machida, Hb Vexin, Hb Corbeil, Hb Ottawa (Siam) #, Hb Fort de France #, Hb S, Hb G-Makassar, Hb Montgomery #, Hb G-Copenhagen, Hb S-Antilles, Hb Handsworth #, Hb Lavagna, Hb Poissy #, Hb Hamadan, Hb Belfast, Hb Russ (alpha 1) #, Hb Russ (alpha 2) #, Hb Evanston, Hb Stanleyville-II # !!, Hb Cocody !!, Hb Reims # !!, variant de Hb A2 "Tokoname" #, variant de Hb A2 "Wayne" (pic 1) #, variant de Hb A2 "Pisa" #, variant de Hb A2 "J-Oxford" #, variant de Hb A2 "Lombard" #, variant de Hb A2 "Tatras" #, variant de Hb A2 "J-Cape Town" (alpha 2) #, variant de Hb A2 "Thionville" #, variant de Hb A2 "J-Cape Town" (alpha 1) #, variant de Hb A2 "Cemenelum" #, variant de Hb A2 "Nikaia" #, variant de Hb A2 "Hopkins-II" (alpha 1) #, variant de Hb A2 "Jackson" (alpha 1) #, variant de Hb A2 "Hopkins-II" (alpha 2) #, variant de Hb A2 "Singapore" # !!, Hb O-Arab dégradée</p> <p><i>Hb Arya # !!, Hb Kenya (HPFH-7), Hb Hasharon (Sinai) #, Hb Dhofar (Yukuhashi), Hb Shimonoseki (Hikoshima) #, Hb O-Indonesia (Buginese-X) #, Hb Machida, Hb Vexin, Hb Corbeil, Hb Ottawa (Siam) #, Hb Fort de France #, Hb S, Hb G-Makassar, Hb Montgomery #, Hb G-Copenhagen, Hb S-Antilles, Hb Handsworth #, Hb Lavagna, Hb Poissy #, Hb Hamadan, Hb Belfast, Hb Russ (alpha 1) #, Hb Russ (alpha 2) #, Hb Evanston, Hb Stanleyville-II # !!, Hb Cocody !!, Hb Reims # !!, "Tokoname" Hb A2 variant #, "Wayne" Hb A2 variant (peak 1) #, "Pisa" Hb A2 variant #, "J-Oxford" Hb A2 variant #, "Lombard" Hb A2 variant #, "Tatras" Hb A2 variant #, "J-Cape Town" Hb A2 variant (alpha 2) #, "Thionville" Hb A2 variant #, "J-Cape Town" Hb A2 variant (alpha 1) #, "Cemenelum" Hb A2 variant #, "Nikaia" Hb A2 variant #, "Hopkins-II" Hb A2 variant (alpha 1) #, "Jackson" Hb A2 variant (alpha 1) #, "Hopkins-II" Hb A2 variant (alpha 2) #, "Singapore" Hb A2 variant # !!, denatured Hb O-Arab</i></p>

TABLEAU / TABLE

CAPI 3 HEMOGLOBIN(E) : VARIANTS POTENTIELS PRÉSENTS DANS CHAQUE ZONE
POTENTIAL VARIANTS LOCATED IN EACH ZONE

Zone	Hémoglobines / Hemoglobins (Hb)
Z(D)	Hb Memphis # !!, Hb G-Audhail # !!, Hb G-Szuhu (Gifu) !!, Hb Leiden !!, Hb Beograd (D-Camperdown), Hb Muravera, Hb D-Bushman, Hb Gavello, Hb Sogn, Hb Matsue-Oki #, Hb Osu Christiansborg, Hb D-Punjab (D-Los Angeles), Hb Watts #, Hb A2-Coburg, Hb G-Waimanalo (Aida) #, Hb Q-India #, Hb Muskegon, Hb D-Ibadan, Hb Buenos Aires (pic mineur) #, Hb Lepore-BW, Hb Q-Iran #, Hb Akron, Hb Summer Hill, Hb G-Philadelphia #, Hb Karlskoga, Hb D-Ouled Rabah, Hb Aichi, Hb Oleander, Hb Yaizu, Hb Kenitra, Hb San Antonio #, Hb Aalborg, Hb Al-Hammadi Riyadh, Hb Ocho Rios, Hb Rocky Mountain (Paddington), Hb Lepore-Hollandia, Hb Quin-Hai, Hb Fort Worth, Hb Mizushi #, Hb G-Honolulu (G-Chinese), Hb Redondo (Isehara), Hb Lepore-Baltimore, Hb G-Ferrara, Hb Djelfa (pic 2) #, Hb G-Hsi-Tsou, Hb Hackney (Xu Chang), Hb Rothschild, Hb Spanish Town #, Hb Korle-Bu (G-Accra), Hb Khartoum, Hb Moabit, Hb Mobile, Hb Köln (Ube-1) #, Hb G-Norfolk #, Hb Ube-4 #, Hb Maputo, Hb Etobicoke #, Hb D-Iran, Hb Caribbean, Hb Okaloosa, Hb St. Luke's #, Hb G-Taipei, Hb G-Coushatta (G-Saskatoon), Hb Winnipeg #, Hb Canuts [A2], Hb Inkster #, Hb Zürich, Hb G-Pest #, Hb P-Galveston, Hb Queens (Ogi) #, Hb Canuts, Hb Aubenas, Hb Setif #, Hb P-Nilotic, Hb G-Galveston (G-Port Arthur), Hb Sunshine Seth # !!, Hb King's Mill !!, Hb Henri Mondor !!, Hb Titusville # !!, variant de Hb A2 "J-Sardegna" # !!, variant de Hb A2 "Suresnes" # !!, variant de Hb A2 "J-Meerut" (alpha 2) #, variant de Hb A2 "J-Broussais" (alpha 2) #, variant de Hb A2 "J-Rajappen" #, variant de Hb A2 "J-Anatolia" #, variant de Hb A2 "J-Meerut" (alpha 1) #, variant de Hb A2 "Ube-2" #, variant de Hb A2 "J-Broussais" (alpha 1) #, variant de Hb A2 "J-Abidjan" #, variant de Hb A2 "J-Toronto" (alpha 1) #, variant de Hb A2 "Mexico" (alpha 2) #, variant de Hb A2 "Thailand" #, variant de Hb A2 "Mexico" (alpha 1) #, variant de Hb A2 "J-Tongariki" #, variant de Hb A2 "Belliard" #, variant de Hb A2 "Neuilly-sur-Marne" #, variant de Hb A2 "J-Wenchang-Wuming" #, variant de Hb A2 "J-Paris-I" (alpha 2) #, variant de Hb A2 "J-Habana" #, variant de Hb A2 "J-Paris-I" (alpha 1) #, Hb E dégradée
	<i>Hb Memphis # !!, Hb G-Audhail # !!, Hb G-Szuhu (Gifu) !!, Hb Leiden !!, Hb Beograd (D-Camperdown), Hb Muravera, Hb D-Bushman, Hb Gavello, Hb Sogn, Hb Matsue-Oki #, Hb Osu Christiansborg, Hb D-Punjab (D-Los Angeles), Hb Watts #, Hb A2-Coburg, Hb G-Waimanalo (Aida) #, Hb Q-India #, Hb Muskegon, Hb D-Ibadan, Hb Buenos Aires (minor peak) #, Hb Lepore-BW, Hb Q-Iran #, Hb Akron, Hb Summer Hill, Hb G-Philadelphia #, Hb Karlskoga, Hb D-Ouled Rabah, Hb Aichi, Hb Oleander, Hb Yaizu, Hb Kenitra, Hb San Antonio #, Hb Aalborg, Hb Al-Hammadi Riyadh, Hb Ocho Rios, Hb Rocky Mountain (Paddington), Hb Lepore-Hollandia, Hb Quin-Hai, Hb Fort Worth, Hb Mizushi #, Hb G-Honolulu (G-Chinese), Hb Redondo (Isehara), Hb Lepore-Baltimore, Hb G-Ferrara, Hb Djelfa (peak 2) #, Hb G-Hsi-Tsou, Hb Hackney (Xu Chang), Hb Rothschild, Hb Spanish Town #, Hb Korle-Bu (G-Accra), Hb Khartoum, Hb Moabit, Hb Mobile, Hb Köln (Ube-1) #, Hb G-Norfolk #, Hb Ube-4 #, Hb Maputo, Hb Etobicoke #, Hb D-Iran, Hb Caribbean, Hb Okaloosa, Hb St. Luke's #, Hb G-Taipei, Hb G-Coushatta (G-Saskatoon), Hb Winnipeg #, Hb Canuts [A2], Hb Inkster #, Hb Zürich, Hb G-Pest #, Hb P-Galveston, Hb Queens (Ogi) #, Hb Canuts, Hb Aubenas, Hb Setif #, Hb P-Nilotic, Hb G-Galveston (G-Port Arthur), Hb Sunshine Seth # !!, Hb King's Mill !!, Hb Henri Mondor !!, Hb Titusville # !!, "J-Sardegna" Hb A2 variant # !!, "Suresnes" Hb A2 variant # !!, "J-Meerut" Hb A2 variant (alpha 2) #, "J-Broussais" Hb A2 variant (alpha 2) #, "J-Rajappen" Hb A2 variant #, "J-Anatolia" Hb A2 variant #, "J-Meerut" Hb A2 variant (alpha 1) #, "Ube-2" Hb A2 variant #, "J-Broussais" Hb A2 variant (alpha 1) #, "J-Abidjan" Hb A2 variant #, "J-Toronto" Hb A2 variant (alpha 1) #, "Mexico" Hb A2 variant (alpha 2) #, "Thailand" Hb A2 variant #, "Mexico" Hb A2 variant (alpha 1) #, "J-Tongariki" Hb A2 variant #, "Belliard" Hb A2 variant #, "Neuilly-sur-Marne" Hb A2 variant #, "J-Wenchang-Wuming" Hb A2 variant #, "J-Paris-I" Hb A2 variant (alpha 2) #, "J-Habana" Hb A2 variant #, "J-Paris-I" Hb A2 variant (alpha 1) #, denatured Hb E</i>
Z(F)	Hb F, Hb Willamette !!, Hb Hoshida (Chaya) !!, Hb Languidic, Hb Chiapas, Hb P-India, Hb Tamano, Hb Sunnybrook, Hb Park Ridge #, Hb Delzicht #, Hb Atago, Hb Deer Lodge, Hb Alabama, Hb Chapel Hill #, Hb Bunbury, Hb Tak, Hb Q-Thailand (G-Taichung) #, Hb Sabine #, Hb Bassett #, Hb Boyle Heights, Hb Les Lilas #, Hb Rampa, Hb Haaglanden, Hb G-Georgia, Hb Barcelona, Hb G-San José, Hb Denmark Hill, Hb Pôrto Alegre, Hb F-Sardinia, Hb Geldrop Santa Anna, Hb Ta-Li, Hb Chongqing, Hb Richmond, Hb Hirose, Hb Abruzzo, Hb Boumerdes #, Hb British Columbia, Hb Kansas, Hb Tarrant #, Hb Verdun #, Hb Swan River #, Hb Attleboro, Hb Sawara, Hb Burke, Hb Dunn #, Hb Manitoba-I #, Hb Manitoba-II #, Hb Sassari #, Hb Hazebrouck !!, Hb Port Phillip !!, Hb Vanderbilt !!, variant de Hb A2 "J-Rovigo" # !!, variant de Hb A2 "Wayne" (pic 2) # !!, Hb S dégradée, Hb D-Punjab dégradée
	<i>Hb F, Hb Willamette !!, Hb Hoshida (Chaya) !!, Hb Languidic, Hb Chiapas, Hb P-India, Hb Tamano, Hb Sunnybrook, Hb Park Ridge #, Hb Delzicht #, Hb Atago, Hb Deer Lodge, Hb Alabama, Hb Chapel Hill #, Hb Bunbury, Hb Tak, Hb Q-Thailand (G-Taichung) #, Hb Sabine #, Hb Bassett #, Hb Boyle Heights, Hb Les Lilas #, Hb Rampa, Hb Haaglanden, Hb G-Georgia, Hb Barcelona, Hb G-San José, Hb Denmark Hill, Hb Pôrto Alegre, Hb F-Sardinia, Hb Geldrop Santa Anna, Hb Ta-Li, Hb Chongqing, Hb Richmond, Hb Hirose, Hb Abruzzo, Hb Boumerdes #, Hb British Columbia, Hb Kansas, Hb Tarrant #, Hb Verdun #, Hb Swan River #, Hb Attleboro, Hb Sawara, Hb Burke, Hb Dunn #, Hb Manitoba-I #, Hb Manitoba-II #, Hb Sassari #, Hb Hazebrouck !!, Hb Port Phillip !!, Hb Vanderbilt !!, "J-Rovigo" Hb A2 variant # !!, "Wayne" Hb A2 variant (peak 2) # !!, denatured Hb S, denatured Hb D-Punjab</i>
Z8	Hb F acétylée, Hb Grifton !!, Hb Lansing !!, Hb Hinsdale !!, Hb Ypsilanti (Ypsi - pic 1) # !!, Hb Auckland !!, Hb Roanne, Hb Southampton (Casper), Hb Yakima, Hb Saint Mandé, Hb Alberta, Hb Bruxelles, Hb Beth Israel, Hb Val de Marne (Footscray) #, Hb Kempsey (Leslie), Hb Atlanta, Hb Chemilly, Hb S-Clichy, Hb Sarrebourg, Hb Ypsilanti (Ypsi - pic 2) #, Hb Charolles #, Hb Athens-GA (Waco), Hb Debrousse, Hb Köln (Ube-1) #, Hb Aubagne, Hb Rainier
	Acetylated Hb F, Hb Grifton !!, Hb Lansing !!, Hb Hinsdale !!, Hb Ypsilanti (Ypsi - peak 1) # !!, Hb Auckland !!, Hb Roanne, Hb Southampton (Casper), Hb Yakima, Hb Saint Mandé, Hb Alberta, Hb Bruxelles, Hb Beth Israel, Hb Val de Marne (Footscray) #, Hb Kempsey, Hb Shelby (Leslie), Hb Atlanta, Hb Chemilly, Hb S-Clichy, Hb Sarrebourg, Hb Ypsilanti (Ypsi - peak 2) #, Hb Charolles #, Hb Athens-GA (Waco), Hb Debrousse, Hb Köln (Ube-1) #, Hb Aubagne, Hb Rainier

TABLEAU / TABLE

CAPI 3 HEMOGLOBIN(E) : VARIANTS POTENTIELS PRÉSENTS DANS CHAQUE ZONE
POTENTIAL VARIANTS LOCATED IN EACH ZONE

Zone	Hémoglobines / Hemoglobins (Hb)
	<p>Hb A, Hb Presbyterian *, Hb Roubaix (Poland) * #, Hb Silver Springs *, Hb El Escorial * #, Hb Dallas * #, Hb Phnom Penh *, Hb La Coruna *, Hb Bougardiery-Mali *, Hb Saint Nazaire *, Hb Barika * #, Hb Allentown *, Hb Allison Park *, Hb Alpertown *, Hb Aلدorf *, Hb Alzette *, Hb Anamosa *, Hb Antibes-Juan-Les-Pins *, Hb Arta (pic majeur) * #, Hb Aurillac *, Hb Austin *, Hb Aylesbury *, Hb Aztec *, Hb Bass Hill *, Hb Beirut *, Hb Belleville *, Hb Belluno *, Hb Bethesda *, Hb Bibba *, Hb Bladensburg *, Hb Bogné *, Hb Bonn *, Hb Brem-sur-Mer *, Hb Brest *, Hb Brigham *, Hb Brisbane (Great Lakes) *, Hb Broomhill *, Hb Brugg *, Hb Buenos Aires (Bryn Mawr - pic majeur) * #, Hb Buffalo (Reeuwijk) *, Hb Bushwick *, Hb Caen *, Hb Calvino *, Hb Cardarelli *, Hb Cheverly *, Hb Chicago *, Hb City of Hope *, Hb Coimbra (Ingelheim) *, Hb Columbia Missouri *, Hb Conakry *, Hb Cowltown *, Hb Crete *, Hb Dapu *, Hb Den Haag *, Hb Denver *, Hb Dhaka *, Hb Dhonburi (Neapolis) *, Hb Djelfa (pic 1) * #, Hb Ecuador *, Hb Evans *, Hb Flurlingen *, Hb Fontainebleau *, Hb Frankfurt *, Hb Fukuoka *, Hb Fukuyama *, Hb Geisinger *, Hb Genova (Hyogo) *, Hb Godavari *, Hb Gorwihl (Hinchingsbrooke) *, Hb Gouda *, Hb Grange Blanche *, Hb Groene Hart (Bernalda) *, Hb Grove City *, Hb Guanajuato *, Hb Haelen *, Hb Hamilton *, Hb Hammersmith (Chiba) *, Hb Heathrow *, Hb Hekinan *, Hb Hershey *, Hb Hyden *, Hb Ingleswood *, Hb Iowa *, Hb Iraq-Halabja *, Hb Jabalpur *, Hb Jeddah *, Hb Johnstown *, Hb Kaiser West End *, Hb Kansas City *, Hb King Ecgbert *, Hb Knossos *, Hb Kokomo *, Hb Kosovo *, Hb La Desirade *, Hb Le Lamentin *, Hb Les Andelys *, Hb Linköping (Meilahti) *, Hb Lisbon *, Hb Little Rock *, Hb Louisville (Bucaresti) *, Hb Lulu Island *, Hb Lyon-Bron *, Hb M-Boston (M-Osaka) *, Hb M-Saskatoon (pic majeur) * #, Hb McKees Rocks *, Hb Malay *, Hb Malmö *, Hb Marijampolė *, Hb Marseille (Long Island) *, Hb Matsudo *, Hb Milledgeville *, Hb Minneapolis Laos *, Hb Mizuho * #, Hb Moriguchi *, Hb Mosella *, Hb Nakhon Ratchasima (Aberystwyth) *, Hb Nantes *, Hb Niguarda *, Hb Noko *, Hb Novara *, Hb Okayama *, Hb Oloupona *, Hb Olympia *, Hb Owari *, Hb Ozieri *, Hb Parma *, Hb Part-Dieu *, Hb Perth (Abraham Lincoln) *, Hb Petit Bourg *, Hb Pierre-Bénite *, Hb Pittsburgh *, Hb Pohpei * #, Hb Port Huron *, Hb Potomac *, Hb Pressath *, Hb Princes Risborough *, Hb Puttelange *, Hb Raleigh *, Hb Ramona *, Hb Ravenscourt Park *, Hb Regina *, Hb Rhode Island (Southwark) *, Hb Riccarton *, Hb Rio Claro *, Hb Rotterdam *, Hb Rouen (Ethiopia) *, Hb Saclay *, Hb Saint-Clair *, Hb Saint-Jacques *, Hb St Joseph's *, Hb Saint-Marcellin *, Hb Saki *, Hb San Bruno *, Hb San Diego *, Hb San Martin *, Hb Santa Barnabas (Croxley Green) *, Hb Santa Juana (Serres) *, Hb Savannah *, Hb Saveh *, Hb Sendagi (Warsaw) *, Hb Sheffield *, Hb Sittia *, Hb Sodertalje *, Hb South Florida *, Hb South Milwaukee *, Hb South Yorkshire *, Hb Sydney *, Hb Taradale (Middlesbrough) *, Hb Taybe *, Hb Templeuve *, Hb Torino *, Hb Toulon *, Hb Twin Peaks *, Hb Ty Gard *, Hb Tyne *, Hb Utrecht *, Hb Uzes *, Hb Valletta *, Hb Valme *, Hb Venetia *, Hb Verona *, Hb Vientiane (Grey Lynn) *, Hb Vila Real *, Hb Villejuif *, Hb Villeparisis *, Hb Villeurbanne *, Hb Volga (Drenthe) *, Hb Voorhees *, Hb Washtenaw *, Hb Waterland *, Hb Weesp *, Hb Wembley *, Hb Westmead *, Hb Wiangpapao *, Hb William-Harvey *, Hb Wood *, Hb Worthing *, Hb Yaounde (Mataro) *, Hb Zoetermeer *, Hb Sinai-Baltimore *, Hb M-Milwaukeee-I *, Hb Melusine * #, Hb Pitie-Salpetriere *, Hb Syracuse *, Hb Hounslow, Hb Fort Dodge, Hb Old Dominion (OD/BUt), Hb Camperdown, Hb Duarte !!, Hb Jura (Bamako) # !!</p>
Z(A)	<p><i>Hb A, Hb Presbyterian * #, Hb Roubaix (Poland) * #, Hb Silver Springs *, Hb El Escorial * #, Hb Dallas * #, Hb Phnom Penh *, Hb La Coruna *, Hb Bougardiery-Mali *, Hb Saint Nazaire *, Hb Barika * #, Hb Allentown *, Hb Allison Park *, Hb Alpertown *, Hb Aلدorf *, Hb Alzette *, Hb Anamosa *, Hb Antibes-Juan-Les-Pins *, Hb Arta (main peak) * #, Hb Aurillac *, Hb Austin *, Hb Aylesbury *, Hb Aztec *, Hb Bass Hill *, Hb Beirut *, Hb Belleville *, Hb Belluno *, Hb Bethesda *, Hb Bibba *, Hb Bladensburg *, Hb Bogné *, Hb Bonn *, Hb Brem-sur-Mer *, Hb Brest *, Hb Brigham *, Hb Brisbane (Great Lakes) *, Hb Broomhill *, Hb Brugg *, Hb Buenos Aires (Bryn Mawr, major peak) * #, Hb Buffalo (Reeuwijk) *, Hb Bushwick *, Hb Caen *, Hb Calvino *, Hb Cardarelli *, Hb Cheverly *, Hb Chicago *, Hb City of Hope *, Hb Coimbra (Ingelheim) *, Hb Columbia Missouri *, Hb Conakry *, Hb Cowltown *, Hb Crete *, Hb Dapu *, Hb Den Haag *, Hb Denver *, Hb Dhaka *, Hb Dhonburi (Neapolis) *, Hb Djelfa (peak 1) * #, Hb Ecuador *, Hb Evans *, Hb Flurlingen *, Hb Fontainebleau *, Hb Frankfurt *, Hb Fukuoka *, Hb Fukuyama *, Hb Geisinger *, Hb Genova (Hyogo) *, Hb Godavari *, Hb Gorwihl (Hinchingsbrooke) *, Hb Gouda *, Hb Grange Blanche *, Hb Groene Hart (Bernalda) *, Hb Grove City *, Hb Guanajuato *, Hb Haelen *, Hb Hamilton *, Hb Hammersmith (Chiba) *, Hb Heathrow *, Hb Hekinan *, Hb Hershey *, Hb Hyden *, Hb Ingleswood *, Hb Iowa *, Hb Iraq-Halabja *, Hb Jabalpur *, Hb Jeddah *, Hb Johnstown *, Hb Kaiser West End *, Hb Kansas City *, Hb King Ecgbert *, Hb Knossos *, Hb Kokomo *, Hb Kosovo *, Hb La Desirade *, Hb Le Lamentin *, Hb Les Andelys *, Hb Linköping (Meilahti) *, Hb Lisbon *, Hb Little Rock *, Hb Louisville (Bucaresti) *, Hb Lulu Island *, Hb Lyon-Bron *, Hb M-Boston (M-Osaka) *, Hb M-Saskatoon (main peak) * #, Hb McKees Rocks *, Hb Malay *, Hb Malmö *, Hb Marijampolė *, Hb Marseille (Long Island) *, Hb Matsudo *, Hb Milledgeville *, Hb Minneapolis Laos *, Hb Mizuho * #, Hb Moriguchi *, Hb Mosella *, Hb Nakhon Ratchasima (Aberystwyth) *, Hb Nantes *, Hb Niguarda *, Hb Noko *, Hb Novara *, Hb Okayama *, Hb Oloupona *, Hb Olympia *, Hb Owari *, Hb Ozieri *, Hb Parma *, Hb Part-Dieu *, Hb Perth (Abraham Lincoln) *, Hb Petit Bourg *, Hb Pierre-Bénite *, Hb Pittsburgh *, Hb Pohpei * #, Hb Port Huron *, Hb Potomac *, Hb Pressath *, Hb Princes Risborough *, Hb Puttelange *, Hb Raleigh *, Hb Ramona *, Hb Ravenscourt Park *, Hb Regina *, Hb Rhode Island (Southwark) *, Hb Riccarton *, Hb Rio Claro *, Hb Rotterdam *, Hb Rouen (Ethiopia) *, Hb Saclay *, Hb Saint-Clair *, Hb Saint-Jacques *, Hb St Joseph's *, Hb Saint-Marcellin *, Hb Saki *, Hb San Bruno *, Hb San Diego *, Hb San Martin *, Hb Santa Barnabas (Croxley Green) *, Hb Santa Juana (Serres) *, Hb Savannah *, Hb Saveh *, Hb Sendagi (Warsaw) *, Hb Sheffield *, Hb Sittia *, Hb Sodertalje *, Hb South Florida *, Hb South Milwaukee *, Hb South Yorkshire *, Hb Sydney * #, Hb Taradale (Middlesbrough) *, Hb Taybe *, Hb Templeuve *, Hb Torino *, Hb Toulon *, Hb Twin Peaks *, Hb Ty Gard *, Hb Tyne *, Hb Utrecht *, Hb Uzes *, Hb Valletta *, Hb Valme *, Hb Venetia *, Hb Verona *, Hb Vientiane (Grey Lynn) *, Hb Vila Real *, Hb Villejuif *, Hb Villeparisis *, Hb Villeurbanne *, Hb Volga (Drenthe) *, Hb Voorhees *, Hb Washtenaw *, Hb Waterland *, Hb Weesp *, Hb Wembley *, Hb Westmead *, Hb Wiangpapao *, Hb William-Harvey *, Hb Wood *, Hb Worthing *, Hb Yaounde (Mataro) *, Hb Zoetermeer *, Hb Sinai-Baltimore *, Hb M-Milwaukeee-I *, Hb Melusine * #, Hb Pitie-Salpetriere *, Hb Syracuse *, Hb Hounslow, Hb Fort Dodge, Hb Old Dominion (OD/BUt), Hb Camperdown, Hb Duarte !!, Hb Jura (Bamako) # !!</i></p>

TABLEAU / TABLE

CAPI 3 HEMOGLOBIN(E) : VARIANTS POTENTIELS PRÉSENTS DANS CHAQUE ZONE
POTENTIAL VARIANTS LOCATED IN EACH ZONE

Zone	Hémoglobines / Hemoglobins (Hb)
Z10	Hb Stockholm !!, Hb Créteil, Hb Nouakchott #, Hb M-Iwate (M-Kankakee) #, Hb Wayne (pic 1) #, Hb Complutense, Hb Camden (Tokuchi), Hb Hope
	<i>Hb Stockholm !!, Hb Créteil, Hb Nouakchott #, Hb M-Iwate (M-Kankakee) #, Hb Wayne (peak 1) #, Hb Complutense, Hb Camden (Tokuchi), Hb Hope</i>
Z11	Hb A dégradée, Hb Vaasa, Hb Tacoma, Hb Providence (pic X-Asn) #, Hb Yamagata, Hb Shepherds Bush, Hb Cook, Hb Corsica, Hb Pisa #, Hb K-Woolwich, Hb Lombard #, Hb J-Guantanamo, Hb Andrew Minneapolis, Hb J-Cape Town (alpha 1) #, Hb Kaohsiung (New York), Hb Fannin-Lubbock I, Hb Saint Claude #, Hb Thionville #, Hb Jackson (alpha 2) #, Hb J-Cape Town (alpha 2) #, Hb Strasbourg, Hb Osler (Fort Gordon), Hb Helsinki, Hb Doha, Hb Linwood, Hb J-Auckland, Hb Nancy, Hb Chesapeake, Hb Himeji, Hb Singapore #, Hb Jackson (alpha 1) #, Hb Cemenelum # !!, Hb Tatras # !!, variant de Hb A2 "I (I-Texas)" #
	<i>Denatured Hb A, Hb Vaasa, Hb Tacoma, Hb Providence (X-Asn peak) #, Hb Yamagata, Hb Shepherds Bush, Hb Cook, Hb Corsica, Hb Pisa #, Hb K-Woolwich, Hb Lombard #, Hb J-Guantanamo, Hb Andrew Minneapolis, Hb J-Cape Town (alpha 1) #, Hb Kaohsiung (New York), Hb Fannin-Lubbock I, Hb Saint Claude #, Hb Thionville #, Hb Jackson (alpha 2) #, Hb J-Cape Town (alpha 2) #, Hb Strasbourg, Hb Osler (Fort Gordon), Hb Helsinki, Hb Doha, Hb Linwood, Hb J-Auckland, Hb Nancy, Hb Chesapeake, Hb Himeji, Hb Singapore #, Hb Jackson (alpha 1) #, Hb Cemenelum # !!, Hb Tatras # !!, "I (I-Texas)" Hb A2 variant #</i>
Z12	Hb Bart, Hb Nikaia # !!, Hb Tokoname # !!, Hb J-Cubuquui, Hb Hopkins-II (alpha 1) #, Hb J-Calabria (J-Bari), Hb J-Camagüey, Hb J-Tongariki #, Hb Wayne (pic 2) #, Hb J-Meerut (J-Birmingham - alpha 1) #, Hb Hopkins-II (alpha 2) #, Hb Zaïre, Hb J-Meerut (J-Birmingham - alpha 2) #, Hb Trollhättan, Hb Pyrgos (Mizunami), Hb Providence (pic X-Asp) #, Hb Suresnes #, Hb J-Broussais (Tagawa-I - alpha 2) #, Hb Grady (Dakar - alpha 2), Hb Grady (Dakar - alpha 1), Hb Legnano, Hb Hikari, Hb J-Rajappen #, Hb J-Anatolia #, Hb J-Broussais (Tagawa-I - alpha 1) #, Hb J-Chicago, Hb J-Sardagna #, Hb J-Toronto (alpha 1) #, Hb J-Cordoba, Hb J-Meinung (J-Bangkok), Hb Ube-2 #, Hb Dagestan, Hb J-Cambridge (Rambam), Hb Hofu, Hb J-Abidjan #, Hb Ulm, Hb Belliard #, Hb J-Iran, Hb Riyadh (Karatsu), Hb Mexico (J-Paris-I - alpha 1) #, Hb Mexico (J-Paris-II - alpha 2) #, Hb Neuilly-sur-Marne #, Hb Pontoise (J-Pontoise), Hb Ankara, Hb J-Buda, Hb J-Medellin, Hb J-Paris-I (J-Aljezur - alpha 1) #, Hb Thailand #, Hb J-Habana #, Hb J-Baltimore (N-New Haven), Hb J-Wenchang-Wuming (Anantharaj) #, Hb J-Paris-I (J-Aljezur - alpha 2) #, Hb Beijing, Hb J-Oxford (I-Interlaken) #, Hb K-Ibadan !!
	<i>Hb Bart, Hb Nikaia # !!, Hb Tokoname # !!, Hb J-Cubuquui, Hb Hopkins-II (alpha 1) #, Hb J-Calabria (J-Bari), Hb J-Camagüey, Hb J-Tongariki #, Hb Wayne (peak 2) #, Hb J-Meerut (J-Birmingham - alpha 1) #, Hb Hopkins-II (alpha 2) #, Hb Zaïre, Hb J-Meerut (J-Birmingham - alpha 2) #, Hb Trollhättan, Hb Pyrgos (Mizunami), Hb Providence (X-Asp peak) #, Hb Suresnes #, Hb J-Broussais (Tagawa-I - alpha 2) #, Hb Grady (Dakar - alpha 2), Hb Grady (Dakar - alpha 1), Hb Legnano, Hb Hikari, Hb J-Rajappen #, Hb J-Anatolia #, Hb J-Broussais (Tagawa-I - alpha 1) #, Hb J-Chicago, Hb J-Sardagna #, Hb J-Toronto (alpha 1) #, Hb J-Cordoba, Hb J-Meinung (J-Bangkok), Hb Ube-2 #, Hb Dagestan, Hb J-Cambridge (Rambam), Hb Hofu, Hb J-Abidjan #, Hb Ulm, Hb Belliard #, Hb J-Iran, Hb Riyadh (Karatsu), Hb Mexico (J-Paris-I - alpha 1) #, Hb Mexico (J-Paris-II - alpha 2) #, Hb Neuilly-sur-Marne #, Hb Pontoise (J-Pontoise), Hb Ankara, Hb J-Buda, Hb J-Medellin, Hb J-Paris-I (J-Aljezur - alpha 1) #, Hb Thailand #, Hb J-Habana #, Hb J-Baltimore (N-New Haven), Hb J-Wenchang-Wuming (Anantharaj) #, Hb J-Paris-I (J-Aljezur - alpha 2) #, Hb Beijing, Hb J-Oxford (I-Interlaken) #, Hb K-Ibadan !!</i>
Z13	Hb Al-Ain Abu Dhabi, Hb J-Europa, Hb N-Baltimore (Hopkins-I), Hb J-Rovigo #, Hb J-Lome, Hb Arta (pic mineur) #, Hb J-Norfolk (Kagoshima), Hb Nigeria, Hb J-Kaohsiung (J-Honolulu)
	<i>Hb Al-Ain Abu Dhabi, Hb J-Europa, Hb N-Baltimore (Hopkins-I), Hb J-Rovigo #, Hb J-Lome, Hb Arta (minor peak) #, Hb J-Norfolk (Kagoshima), Hb Nigeria, Hb J-Kaohsiung (J-Honolulu)</i>
Z14	Hb N-Seattle, Hb J-Tashkuergan
	<i>Hb N-Seattle, Hb J-Tashkuergan</i>
Z15	Hb H, Hb I-Toulouse !!, Hb Sudbury, Hb Kurosaki (alpha 1), Poly A (A->G); AATAAA->AATAAG of the alpha2 gene alpha-Thal-2, Hb Kurosaki (alpha 2), Hb F-Emirates, Hb N-Timone, Hb I (I-Texas, I-Philadelphia) #, Hb Shaare Zedek
	<i>Hb H, Hb I-Toulouse !!, Hb Sudbury, Hb Kurosaki (alpha 1), Poly A (A->G); AATAAA->AATAAG of the alpha2 gene alpha-Thal-2, Hb Kurosaki (alpha 2), Hb F-Emirates, Hb N-Timone, Hb I (I-Texas, I-Philadelphia) #, Hb Shaare Zedek</i>

* Pic non ou peu visible car co-migrant de la fraction normale
Hidden or partially hidden peak due to similar migration time to normal fraction

Variant avec plusieurs fractions affichées (variant de la chaîne alpha ou variant instable ...)
Variant with several fractions displayed (alpha-chain or unstable variant ...)

!! Pic en bordure de zone (risque de saut de zone)
Peak in zone boundary (risk of zone shift)

Rappel : dans chaque zone les variants Hb sont listés selon leur temps de migration de droite vers la gauche
Reminder: in each zone Hb variants are sorted according to their migration time from right to left

TABLEAU / TABLE

FR : VARIANTS POTENTIELS PRÉSENTS DANS CHAQUE ZONE	Zone	Hémoglobines (Hb)	* Pic peu ou pas visible car migrant conjointement avec la fraction normale
GB : POTENTIAL VARIANTS LOCATED IN EACH ZONE	Zone	Hemoglobins (Hb)	* Hidden or partially hidden peak due to similar migration time to normal fraction
DE : POTENZIELLE VARIANTE IN DEN EINZELNEN ZONEN	Zone	Hämoglobine (Hb)	* Versteckter oder teilweise versteckter Spitzenwert infolge ähnlicher Migrationszeit wie bei der normalen Fraktion
NL : POTENTIELE VARIANTEN IN ELKE ZONE	Zone	Hemoglobinen (Hb)	* Verborgen of gedeeltelijk verborgen pik vanwege migratietijd die vergelijkbaar is met normale fractie
IT : VARIANTI POTENZIALI PRESENTI IN CIASCUNA ZONA	Zona	Emoglobine (Hb)	* Picco invisibile o scarsamente visibile poiché migrante unitamente alla frazione normale
ES : VARIANTES POTENCIALES PRESENTES EN CADA ZONA	Zona	Hemoglobinas (Hb)	* Pico oculto o parcialmente oculto debido a migración conjunta con la fracción normal
PT : VARIANTES POTENCIAIS LOCALIZADAS EM CADA ZONA	Zona	Hemoglobinas (Hb)	* Pico oculto ou parcialmente oculto devido a tempo de migração similar para fração normal
SV : POTENTIALIA VARIANTER BELÄGNA I VARJE ZON	Zon	Hemoglobiner (Hb)	* Dold eller delvis dold topp på grund av liknande migratinstid till normal fraktion
GR : ΔΥΝΗΤΙΚΕΣ ΠΑΡΑΜΕΤΡΕΣ ΠΟΥ ΕΝΤΟΠΙΖΟΝΤΑΙ ΣΕ ΚΑΘΕ ΖΩΝΗ	Ζώνη	Αιμοσφαιρίνες (Hb)	* Απόκρυψη ή μερικώς απόκρυψη κορυφής λόγω παρόμοιου χρόνου ηλεκτροφόρησης σε φυσιολογική κλάση
HR : POTENCIJALNE VARIJANTE LOČRANJE U SVAKOJ ZONI	Zona	Hemoglobini (Hb)	* Vrh je potpuno ili djelomično skriven zbog vremena migracije sličnog s onom za normalnu frakciju
IT : GALIMI VARIANTI, ESANTYI KIEVNIENJE ZONJUE	Zona	Hemoglobinas (Hb)	* Pasklata arba iš dalies paslepta pikšnis, nes migracijos laikas panašus į normalios frakcijos migracijos laiką
PL : POTENCJALNE ODMIANY ZLOKALIZOWANE W KAŻDEJ STREFIE	Strefa	Hemoglobiny (Hb)	* Ukryty lub częściowo ukryty pik ze względu na podobny czas migracji względem frakcji prawidłowej
RO : VARIANTIE POTENȚIALE SITUATE ÎN FIECARE ZONĂ	Zonă	Hemoglobine (Hb)	* Vârf ascuns sau parțial ascuns datorită timpului de migrare asemănător cu fracția normală
CS : POTENCIJALNE VARIJANTE KOJE SE NALAZE U SVAKOJ ZONI	Zona	Hemoglobini (Hb)	* Sakriveni ili djelimično sakriveni vrh zbog sličnog vremena migracije u normalnu frakciju
HU : LEHETSÉGESES VARIÁNSOK AZ EGYES ZÓNÁKBAN	Zóna	Hemoglobinek (Hb)	* A normál frakciónalhoz hasonló migrációs idő miatt rejtett vagy részben rejtett csúcs
TR : HER BİR BÖLGEDE YER ALAN VARYANTLAR	Bölge	Hemoglobinler (Hb)	* Normal fraksiyona benzer migrasyon süresi nedeniyle gizli veya kısmen gizli pik
CZ : POTENCIJALNÍ VARIANTY HEMOGLOBINU UMÍSTĚNÉ V KAŽDÉ ZÓNĚ	Zóna	Hemoglobiny (Hb)	* Skrytý nebo částečně skrytý pik v důsledku podobné doby migrace normální frakce
BG : ВЪЗМОЖНИ ВАРИАНТИ, РАЗПОЛОЖЕНИ ВЪВ ВСЯКА ЗОНА	Зона	Хемоглобини (Hb)	* Скрип или частично скрип пик поради време на миграция, подобно на това на нормалната фракция
NO : POTENSIELLE VARIANTER PLESSERT I HVER SONE	Sone	Hemoglobiner (Hb)	* Skjult eller delvis skjult topp på grunn av lignende migreringstid til normal fraksjon
DK : POTENTIELLE VARIANTER I HVER ZONE	Zone	Hæmoglobiner (Hb)	* Skjult eller delvis skjult topp på grund af lignende migratinstid til normal fraktion
CN : 每个区中潜在的变种	区	血红蛋白 (Hb)	* 由于电泳时间与普通组分相似，因此峰被掩盖或部分掩盖
RU : ВОЗМОЖНЫЕ ВАРИАНТЫ, РАСПОЛОЖЕННЫЕ В КАЖДОЙ ЗОНЕ	Зона	Гемоглобин (Hb)	* Скрытый или частично скрытый пик по причине схожего времени миграции в нормальную фракцию
JP : 各ゾーンに位置する潜在的な変異体	ゾーン	ヘモグロビン (Hb)	* 正常フラクシヨンの移行時間と同一であるため、非表示または部分的に非表示のピーク
LV : POTENCIJALIE VARIANTI KATRĀ ZONĀ	Zona	Hemoglobīns (Hb)	* Slēpta vai daļēji slēpta maksimālā vērtība, ko izraisa migrācijas laiks, kas ir līdzīgs ar normālu frakciju
SK : POTENCIJÁLNE VARIANTY, KTORÉ SA NACHÁDZAJÚ V KAŽDEJ ZÓNĚ	Zóna	Hemoglobíny (Hb)	* Skrytá alebo čiastočne skrytá špička v dôsledku podobného času migrácie ako pri normálnej frakcii
EE : POTENTSIJALSED VARIANDID IGAS TSOONIS	Tsoon	Hemoglobiniid (Hb)	* Varjatud või osaliselt varjatud pik normaalise fraktsiooniga sarnase migratsiooniga lõttu
VN : BIẾN THỂ ẨN TỐI MỖI VÙNG	Vùng	Hemoglobin (Hb)	* Đỉnh ẩn hoặc ẩn một phần vì thời gian di chuyển sang mang thời gian thường tương tự

FR : # Variant avec plusieurs fractions affichées (variant de la chaîne alpha ou variant instable ...)			!! Pic en bordure de zone (risque de changement de zone)
GB : # Variant with several fractions displayed (alpha-chain or unstable variant ...)			!! Peak in zone boundary (risk of zone shift)
DE : # Variante mit mehreren angezeigten Fraktionen (Alpha-Kette oder instabile Variante ...)			!! Spitzenwert im Zonenübergangsbereich (Risiko einer Zonenverschiebung)
NL : # Variant met verschillende zichtbare fracties (alfaketen of onstabiele variantie ...)			!! Piek in zonegrens (gevaar voor zonevervorming)
IT : # Variante con più frazioni visualizzate (variante della catena alfa o variante instabile, ecc.)			!! Picco sul confine di zona (rischio di variazione di zona)
ES : # Variante que presenta varias fracciones (variante de cadena alfa o variante inestable ...)			!! Pico en el límite de la zona (riesgo de cambio de zona)
PT : # Variante com várias frações apresentadas (cadeia alfa ou variante instável, etc.)			!! Pico no limite da zona (risco de deslocação de zona)
SV : # Variant med flera fraktioner som visas (alfa-kedja eller instabil variant ...)			!! Topp i zongräns (risk för zonförskjutning)
GR : # Παραλλαγή με εμφάνιση πολλών κλασμάτων (άλλα αλυσίδα ή ασταθής παραλλαγή ...)			!! Κορυφή σε όριο ζώνης (κίνδυνος μετατόπισης ζώνης)
HR : # Varijanta s nekoliko prikazanih frakcija (alfa-lanac ili nestabilna varijanta ...)			!! Vrh u granici zone (opasnost od pomaka zone)
LT : # Varianto su keliomis rodomomis frakcijomis nr. (alfa grandis arba nestabili variantas ...)			!! Viršūnė ant zonos ribos (zonos paslinkimo pavojus)
PL : # odmiany z kilkoma wyświetlonymi frakcjami (alfacuch lub niestabilna niestabilna ...)			!! Pik w granicy strefy (ryzyko przesunięcia strefy)
RO : # varianta cu mai multe fractii afisate (lant alfa sau varianta instabila ...)			!! Vârf în limita zonei (risic de schimbare a zonei)
CS : # Varijanta sa prikazanih niekoľko frakcijs (alfa lanec ili nestabilna varijanta ...)			!! Vrh u hranici zóny (riziko od pomaku zone)
HU : # Több fragmentumot mutatató variáns (alfa láncc vagy instabil variáns ...)			!! Csúcs a zona határánál (zónáeltolódás kockázata)
TR : # Birden çok fraksiyonu görünlüklendiren varyant (alfa zincir veya kararsız varyant ...)			!! Bölge sınırında pik değeri (bölge kayması riski)
CZ : # Varianta s několika zobrazenými frakcemi (alfa řetězec nebo nestálá varianta ...)			!! Pik na hranici zóny (riziko posunutí zóny)
BG : # Появеща се евариант с няколко фракции (алфа-верига или нестабилна евариант ...)			!! Пик на границата на зоната (опасност от преместване на зоната)
NO : # Variant med flere fraksjoner vises (Alpha-kjeden eller ustabil variant ...)			!! Topp i sonegrense (risiko for soneforskyvning)
DK : # Variant med flere fraktioner vist (alfakæde eller ustabil variant ...)			!! Top i zonegrænse (fare for zoneforskyvning)
CN : # 显示了多个区分的变体 (α 链或不稳定变体 ...)			!! 区边界峰位 (有区位移的风险)
RU : # Вариация с отображением нескольких фракций (альфа-цепь или неустойчивая вариация ...)			!! Пик на границе зоны (риск смещения зоны)
JP : # 数個のフラクシヨンを表示される変異体 (α 鎖または不安定な変異体 ...)			!! ゾーン境界でのピーク (ゾーンシフトのリスクがある)
LV : # Tiek parādīts variants ar vairākiem frakcijām (alfa ķēde vai nestabila varianta u.c.)			!! Maksimālā vērtība zonas robežās (zonas maiņas risks)
SK : # Variant s viacermi zobrazenými frakciami (alfa-řetazec alebo nestabilná varianta ...)			!! Špička v hranici zóny (riziko posunutia zóny)
EE : # Kuivatud on mitme fraktsiooniga variant (alfa-alah või eabestabiilne variant ...)			!! Pikk tsooni piiri (tsooni nihke risk)
VN : # Biến cố vài miền hiển thị (biến chuỗi alpha hay biến không ổn định ...)			!! Đỉnh nằm trong ranh giới vùng (nguy cơ thay đổi vùng)

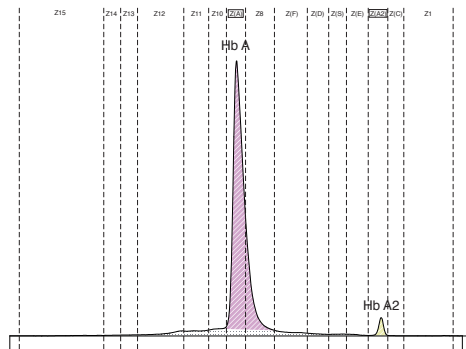
FR : Rappel : Dans chaque zone, les variants Hb sont listés selon leur temps de migration de la droite vers la gauche			
GB : Reminder : In each zone, Hb variants are sorted according to their migration time from the right to the left			
DE : Hinweis : Die Hb-Varianten werden in allen Zonen nach ihrer Migrationszeit von rechts nach links sortiert			
NL : Herinnering : In elke zone worden Hb varianten op basis van hun migratietijd van rechts naar links gesorteerd			
IT : NB : In ciascuna zona, le varianti Hb sono elencate in base al loro tempo di migrazione da destra verso sinistra			
ES : NOTA : En cada zona, las variantes de la Hb son listadas según su tiempo de migración de derecha a izquierda			
PT : Lembrete : Em cada zona, as variantes Hb são ordenadas de acordo com o seu tempo de migração da direita para a esquerda			
SV : PÅminnelse : I varje zon, sorteras Hb-varianter enligt deras migratinstid från höger till vänster			
GR : Υπενήμιση : Σε κάθε ζώνη, οι παραλλαγές Hb ταξινομούνται ανάλογα με τον χρόνο ηλεκτροφόρησης τους από τα δεξιά προς τα αριστερά			
HR : Podsjetnik : Hb varijante razvrstane su u svakoj zoni prema svom vremenu migracije s desne na lijevu stranu			
LT : Pirmename : kiekvienoje zonoje Hb variantai pagal migracijos laiką suriūšomi iš dešinės į kairę.			
PL : Przypomnienie : W każdej strefie odmiany Hb są sortowane według czasu migracji, od prawej do lewej			
RO : Memento : În fiecare zonă, variantele Hb sunt sortate în funcție de timpul de migrare de la dreapta la stânga			
CS : Podsetník : U svakovj zoni, Hb varijante se sortují prema svom vremenu migracijsa desna na levo			
HU : Émlékeztető : A Hb-variánsok mindenkgy zónában a migrációs idejüknek megfelelően rendeződnek jobbról balra			
TR : Hatırlatma : Her bir bölgede, Hb varyantları sağdan sola migrasyon sürelerine göre sımiflandırılır			
CZ : Pripomínka : V každé zóně jsou varianty Hb rozříděny podle své doby migrace zprava doleva			
BG : Напоминание : Hb вариантите във всяка зона се сортират от дясно наляво според времето им на миграция			
NO : Påminnelse : I hver sone, er Hb-varianter sortert i henhold til deres migreringstid fra høyre til venstre			
DK : Påmindelse : I hver zone sorteres Hb-varianter efter deres migratinstid fra højre til venstre			
CN : 提示 : 在每个区中, 根据 Hb 变体从右到左的电泳时间进行排序			
RU : Напоминание : вариации Hb сортируются по времени миграции в каждой зоне справа налево			
JP : 注意 : 各ゾーンにおいて、Hb変異体は泳動時間に従って右から左にソートされ(並べ替えられ)ます			
LV : Atgādinājums! Katrā zonā Hb varianti tiek sakārtoti pēc to migrācijas laika no labās uz kreiso pusi			
SK : Pripomenka : Varianty Hb sú v každej zóne usporiadané sprava doľava podľa času migrácie			
EE : Meelespea : Igas tsoonis sorteatakse Hb variandid vastavalt nende migratsioonilajale paremalt vasakule.			
VN : Xin nhắc lại : Ở mỗi vùng, biến thể Hb được sắp xếp theo thời gian di chuyển từ phải sang trái			

SCHÉMAS / FIGURES

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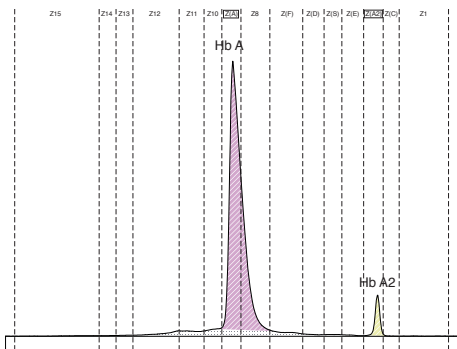
CAP1 3 HEMOGLOBIN(E)
 PROFILS ÉLECTROPHORÉTIQUES - ELECTROPHORETIC PATTERNS

1



Sang normal
 Normal blood sample

2



Sang bêta-thalassémique
 Blood sample with beta-thalassemia

FR : PROFILS ÉLECTROPHORÉTIQUES
 GB : ELECTROPHORETIC PATTERNS
 DE : ELEKTROPHORESEMUSTER
 NL : ELEKTROFORETISCHE PATRONEN
 IT : PROFILI ELETTROFORETICI
 ES : PERFILES ELECTROFORÉTICOS
 PT : PADRÕES ELETTROFORÉTICOS
 SV : ELEKTROFORETISKA MÖNSTER
 GR : ΗΛΕΚΤΡΟΦΟΡΗΤΙΚΑ ΠΡΟΤΥΠΑ
 HR : ELEKTROFORETSKI OBRASCI
 LT : ELEKTROFORĖZĖS ŠABLONAI
 PL : OBRAZY ELEKTROFORETYCZNE
 RO : TIPARE ELECTROFORETICE
 CS : ELEKTROFORETSKI ŠABLONI
 HU : ELEKTROFORETIKUS MINTÁZATOK
 TR : ELEKTROFORETIK PATERNLER
 CZ : ELEKTROFORETIKÉ TYPU
 BG : ELEKTROFORETIЧНИ МОДЕЛИ
 NO : ELEKTROFORETISKE MØNSTER
 DK : ELEKTROFORETISKE MØNSTER
 CN : 电泳图谱
 RU : ЭЛЕКТРОФОРЕТИЧЕСКИЕ ПРОФИЛИ
 JP : 電気泳動パターン
 LV : ELEKTROFORETIŠKIE SPEKTRI
 SK : ELEKTROFORÉŽNE VZORY
 EE : ELEKTROFORETILISED MÜSTRID
 VN : MÔ HÌNH ĐIỆN DI

Sang normal
 Normal blood sample
 Normalblutprobe
 Normaal bloedmonster
 Sangue normale
 Sangre normal
 Amostra de sangue normal
 Normalt blodprov
 Φυσιολογικό δείγμα αίματος
 Normalan uzorak krvi
 Normalus kraujo mėginys
 Próba krwi prawidłowej
 Próba normală de sânge
 Normalan uzorak krvi
 Normál vérminta
 Normal kan numnesi
 Normalni vzorek krve
 Normalna кръвна проба
 Normal blodprobe
 Normal blodprobe
 正常血液样品
 Образец нормальной крови
 正常血液サンプル
 Normāls asins paraugs
 Vzorka normālej krvi
 Normalne vereproov
 Mẫu máu thông thường

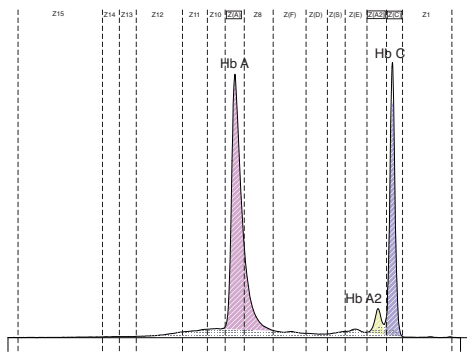
Sang bêta-thalassémique
 Blood sample with beta-thalassemia
 Blutprobe mit Beta-Thalassämie
 Bloedmonster met bètathalassemie
 Sangue beta-talassémico
 Sangre con beta talasemia
 Amostra de sangue con beta-talasemia
 Blodprov med beta-thalassemi
 Δείγμα αίματος με βίητο-θαλασσαιμία
 Uzorak krvi s beta-talassemijom
 Paciento, sergandio beta talasemija, kraujo mėginys
 Próba krwi z beta-talassemią
 Próba de sânge cu beta-talassemie
 Uzorak krvi sa beta-talassemijom
 Bèta-talassémias vérminta
 Beta-talassemi iðeren kan numnesi
 Vzorek krve s beta talasemii
 Кръвна проба с beta-таласемия
 Blodprobe med beta-talassemi
 Blodprobe med beta-talassemi
 β-地中海貧血的血液样品
 Образец крови с бета-талассемией
 βサラセミアの血液サンプル
 Asins paraugs ar beta talasēmiju
 Vzorka krvi s beta-talassēmiu
 Beeta-talassemiiga vereproov
 Mẫu máu có beta-thalassemia

SCHÉMAS / FIGURES

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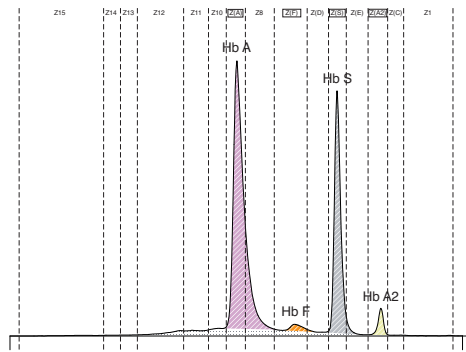
**CAPI 3 HEMOGLOBIN(E)
PROFILS ÉLECTROPHORÉTIQUES - ELECTROPHORETIC PATTERNS**

3



Sang de patient hétérozygote avec variant Hb C
Blood sample from heterozygous patient with Hb C variant

4



Sang de patient hétérozygote avec variant Hb S
Blood sample from heterozygous patient with Hb S variant

- FR : Sang de patient hétérozygote avec variant Hb C
- GB : Blood sample from heterozygous patient with Hb C variant
- DE : Blutprobe eines heterozygoten Patienten mit Hb C-Variante
- NL : Bloedmonster van heterozygote patiënt met Hb C variant
- IT : Sanguè di paziente eterozygote con variante Hb C
- ES : Sangre de paciente heterocigoto con la variante Hb C
- PT : Amostra de sangue de doente heterocigótico com variante Hb C
- SV : Blodprov från heterozygot patient med Hb C-variant
- GR : Δείγμα αίματος από ετερόζυγο ασθενή με παραλλαγή Hb C
- HR : Uzorak krvi heterozigotnog bolesnika s varijantom Hb C
- LT : Heterozigotinio paciento kraujo, kuriame yra Hb C varianto, kraujo mėginys
- PL : Próbk krwi od heterozygotycznego pacjenta z odmianą Hb C
- RO : Probă de sânge de la pacient heterozygot cu varianta Hb C
- CS : Uzorak krvi od heterozygotnog pacijenta sa Hb C varijantom
- HU : Heterozigóta beteg vérmintéjára Hb C variánsal
- TR : Hb C varyantı taşıyan heterozygot hastasına ait kan numunesi
- CZ : Vzorak krve heterozygotního pacienta s variantou Hb C
- BG : Кръвна проба от хетерозиготен пациент с Hb C вариант
- NO : Blodprobe fra heterozygot patient med Hb C-variant
- DK : Blodprobe fra heterozygot patient med Hb C-variant
- CN : 来自 Hb C 杂体杂合患者的血液样品
- RU : Образец крови от гетерозиготного пациента с вариацией Hb C
- JP : Hb C 変異体を含むヘテロ雑合患者からの血液サンプル
- LV : Heterozigota pacienta asins paraugs ar Hb C variantu
- SK : Vzorka krvi od heterozygotného pacienta s variantom Hb C
- EE : Vereproov Hb C variantiga heterosügootselt patsiendilt
- VN : Mẫu máu của bệnh nhân bị bệnh dị hợp tử với biến thể Hb C

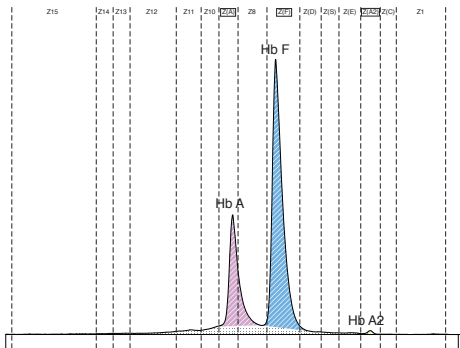
- Sang de patient hétérozygote avec variant Hb S
- Blood sample from heterozygous patient with Hb S variant
- Blutprobe eines heterozygoten Patienten mit Hb S-Variante
- Bloedmonster van heterozygote patiënt met Hb S variant
- Sanguè di paziente eterozygote con variante Hb S
- Sangre de paciente heterocigoto con la variante Hb S
- Amostra de sangue de doente heterocigótico com variante Hb S
- Blodprov från heterozygot patient med Hb S-variant
- Δείγμα αίματος από ετερόζυγο ασθενή με παραλλαγή Hb S
- Uzorak krvi heterozigotnog bolesnika s varijantom Hb S
- Heterozigotinio paciento kraujo, kuriame yra Hb S varianto, mėginys
- Próbka krwi od heterozygotycznego pacjenta z odmianą Hb S
- Probă de sânge de la pacient heterozygot cu varianta Hb S
- Uzorak krvi od heterozygotnog pacijenta sa Hb S varijantom
- Heterozigóta beteg vérmintéjára Hb S variánsal
- Hb S varyantı taşıyan heterozygot hastasına ait kan numunesi
- Vzorak krve heterozygotního pacienta s variantou Hb S
- Кръвна проба от хетерозиготен пациент с Hb S вариант
- Blodprobe fra heterozygot patient med Hb S-variant
- Blodprobe fra heterozygot patient med Hb S-variant
- 来自 Hb S 杂体杂合患者的血液样品
- Образец крови от гетерозиготного пациента с вариацией Hb S
- Hb S 変異体を含むヘテロ雑合患者からの血液サンプル
- Heterozigota pacienta asins paraugs ar Hb S variantu
- Vzorka krvi od heterozygotného pacienta s variantom Hb S
- Vereproov Hb S variantiga heterosügootselt patsiendilt
- Mẫu máu của bệnh nhân bị bệnh dị hợp tử với biến thể Hb S

SCHÉMAS / FIGURES

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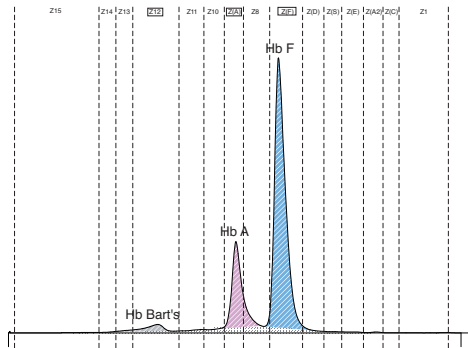
CAPI 3 HEMOGLOBIN(E)
 PROFILS ÉLECTROPHORÉTIQUES - ELECTROPHORETIC PATTERNS

5



Sang normal de bébé (âgé de 3 semaines)
 Normal blood sample from baby (3 weeks old)

6



Sang de bébé avec Hb Bart
 Baby blood sample with Hb Bart's

FR : Sang normal de bébé (âgé de 3 semaines)
 GB : Normal blood sample from baby (3 weeks old)
 DE : Normalblutprobe eines Säuglings (Alter: 3 Wochen)
 NL : Normaal bloedmonster van baby (3 weken oud)
 IT : Sangue normale di neonato (età 3 settimane)
 ES : Sangre normal de bebé (3 semanas de edad)
 PT : Amostra de sangue normal de bebé (3 semanas de idade)
 SV : Normalt blodprov från baby (3 veckor gammal)
 GR : Φυσιολογικό δείγμα αίματος από βρέφος (ηλικίας 3 εβδομάδων)
 HR : Normalan uzorak krvi dojenčeta (u dobi od 3 tjedna)
 LT : Normalus kūdikio (3 savaičių amžiaus) kraujo mėginys
 PL : Próba krwi prawidłowej od niemowlęcia (3-tygodniowego)
 RO : Probă normală de sânge de la bebeluș (trei săptămâni)
 CS : Normální uzorak krvi od bebe (stare 3 neděle)
 HU : 3 hetes csecsemő normál vérmintája
 TR : Bebekten alınan normal kan numunesi (3 haftalık)
 CZ : Normální vzorek krve malého dítěte (věk 3 týdnů)
 BG : Нормална кръвна проба от бебе (на възраст 3 седмици)
 NO : Normal blodprøve fra nyfødt barn (3 uker gammel)
 DK : Normal blodprøve fra baby (3 uger gammel)
 CN : 来自婴儿 (3 个月) 的正常血液样品
 RU : Образец нормальной крови младенца (возраст — 3 недели)
 JP : 新生児 (生後3週) からの正常血液サンプル
 LV : Mazuļa (3 nedēļas vecs) normāls asiņu paraugs
 SK : Vzorka normálnej krvi od novorodenca (vo veku 3 týždňov)
 EE : Normaalne vereproov imikult (3-nädalane)
 VN : Mẫu máu của trẻ sơ sinh (3 tuần tuổi)

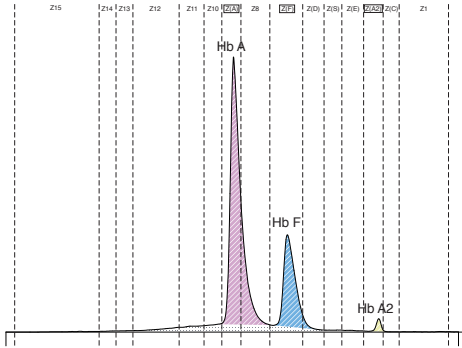
Sang de bébé avec Hb Bart
 Baby blood sample with Hb Bart's
 Blutprobe eines Säuglings mit Hb-Barts
 Bloedmonster van baby met Hb Bart
 Sangue di neonato con Hb Bart
 Sangre de bebé con Hb Bart
 Amostra de sangue de bebé com Hb de Bart
 Blodprov från baby med Hb Bart's
 Δείγμα αίματος βρέφους με Hb Bart's
 Uzorak krvi dojenčeta s Bartovim Hb
 Kūdikio kraujo, kuriame yra Hb Bart's, mėginys
 Próba krwi niemowlęcia z Hb Barta
 Probă de sânge de la bebeluș cu Hb Bart's
 Uzorak krvi bebe sa Hb Barts
 Csecsemő vérmintája Hb Barttal
 Hb Barts taşıyan bebeğin ait kan numunesi
 Vzorek krve malého dítěte s Bartovým Hb
 Кръвна проба от бебе с Hb на Bart
 Blodprøve fra nyfødt barn med Hb Barts
 Babyblodprøve med Hb Bart's
 含 Hb Bart's 的血液样品
 Образец крови младенца с гемоглобином Барта
 Hb(バーツ)を含む新生児の血液サンプル
 Mazuļa asiņu paraugs ar Hb Bart
 Vzorka krvi od novorodenca s Hb Bartovým Hb
 Imiku vereproov Hb Bart'iga
 Mẫu máu của trẻ sơ sinh có Hb Bart's

SCHÉMAS / FIGURES

ABBILDUNGEN - FIGUREN - FIGURE - FIGURAS - BILDER - ΕΙΚΟΝΕΣ - SLIKE - PAVEIKSLAI - RYSUNKI - FIGURI - ÁBRÁK - ŞEKİLLER - OBRAZKY - ФИГУРИ - FIGURER - 插图 - РИСУНКИ - 図 - CIPARI - JOONISED - SƠ ĐỒ

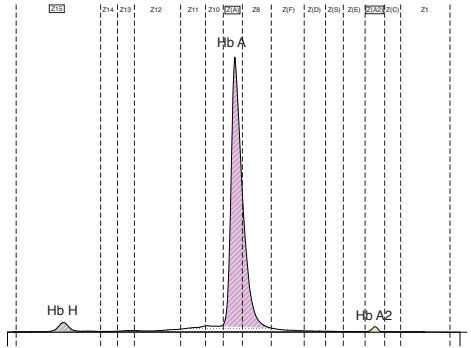
CAPI 3 HEMOGLOBIN(E)
 PROFILS ÉLECTROPHORÉTIQUES - ELECTROPHORETIC PATTERNS

7



Sang avec Hb F élevée (jeune enfant)
 Blood sample with elevated Hb F (young child)

8



Sang avec Hb H
 Blood sample with Hb H

- FR : Sang avec Hb F élevée (jeune enfant)
- GB : Blood sample with elevated Hb F (young child)
- DE : Blutprobe mit erhöhtem Hb F (Kleinkind)
- NL : Bloedmonster met verhoogd Hb F niveau (klein kind)
- IT : Sangue con Hb F alta (bambino)
- ES : Sangre con Hb F elevada (niño de corta edad)
- PT : Amostra de sangue com Hb F elevada (criança pequena)
- SV : Blodprov med förhöjd Hb F (litt barn)
- GR : Δείγμα αίματος με αυξημένο επίπεδο Hb F (μικρό παιδί)
- HR : Uzorak krvi s povišenom vrijednošću Hb F (malo dijete)
- LT : Kraujo, kuriame padidėjęs Hb F kiekis, mėginys (mažas vaikas)
- PL : Próba krwi o podwyższonym stężeniu Hb F (młode dziecko)
- RO : Probă de sânge cu Hb F crescută (copil mic)
- CS : Uzorak krvi sa povišením Hb F (malo dete)
- HU : Vérminta emelkedett Hb F-tel (kisgyermek)
- TR : Yüksek Hb F içeren kan numunesi (genç çocuk)
- CZ : Vzorek krve se zvýšeným Hb F (dítě)
- BG : Кръвна проба с повишен Hb F (малко дете)
- NO : Blodprøve med forhøyet Hb F (små barn)
- DK : Blodprøve med forhøjet Hb F (lille barn)
- CN : Hb F 升高 (幼儿) 的血浆样品
- RU : Образец крови с повышенным уровнем Hb F (ребенок младшего возраста)
- JP : Hb Fが高値の血液サンプル (幼児)
- LV : Asins paraugs ar paaugstinātu Hb F līmeni (mazs bērns)
- SK : Vzorka krvi sa zvýšenou hladinou Hb F (dôčča)
- EE : Vereproov kõrgegenud Hb F-ga (noor laps)
- VN : Mẫu máu có Hb F gia tăng (trẻ nhỏ)

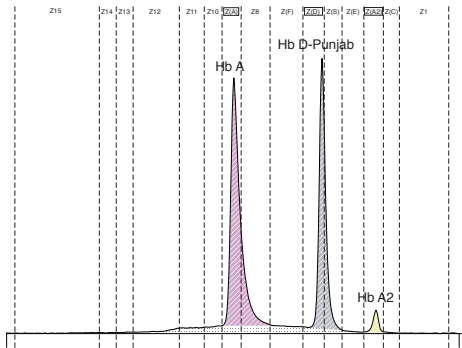
- Sang avec Hb H
- Blood sample with Hb H
- Blutprobe mit Hb H
- Bloedmonster met Hb H
- Sangue con Hb H
- Sangre con Hb H
- Amostra de sangue com Hb H
- Blodprov med Hb H
- Δείγμα αίματος με Hb H
- Uzorak krvi s Hb H
- Kraujo, kuriame yra Hb H, mėginys
- Próba krwi z Hb H
- Probă de sânge cu Hb H
- Uzorak krvi sa Hb H
- Vérminta Hb H-val
- Hb H içeren kan numunesi
- Vzorek krve s Hb H
- Кръвна проба с Hb H
- Blodprøve med Hb H
- Blodprøve med Hb H
- 含 Hb H 的血浆样品
- Образец крови с Hb H
- Hb Hを含む血液サンプル
- Asins paraugs ar Hb H
- Vzorka krvi s Hb H
- Vereproov Hb H-ga
- Mẫu máu có Hb H

SCHÉMAS / FIGURES

ABBILDUNGEN - FIGUREN - FIGURE - FIGURAS - BILDER - EIKONEΣ - SLIKE - PAVEIKSLAI - RYSUNKI - FIGURI - ÁBRÁK - ŞEKİLLER - OBRÁZKY - ΦΙΓΥΡΗ - FIGURER - 插图 - РИСУНКИ - 図 - CIPARI - JOONISED - SƠ ĐỒ

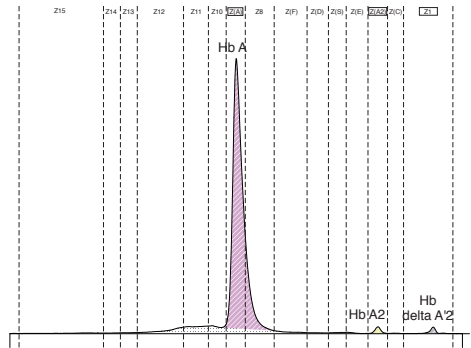
CAPI 3 HEMOGLOBIN(E)
 PROFILS ÉLECTROPHORÉTIQUES - ELECTROPHORETIC PATTERNS

9



Sang de patient hétérozygote avec variant Hb D-Punjab
 Blood sample from heterozygous patient with Hb D-Punjab variant

10



Sang de patient hétérozygote avec variant delta Hb A2
 Blood sample from heterozygous patient with delta Hb A2 variant

- FR : Sang de patient hétérozygote avec variant Hb D-Punjab
- GB : Blood sample from heterozygous patient with Hb D-Punjab variant
- DE : Blutprobe eines heterozygoten Patienten mit Hb D-Punjab-Variante
- NL : Bloedmonster van heterozygote patiënt met Hb D-Punjab variant
- IT : Sangue di paziente eterozygote con variante Hb D-Punjab
- ES : Sangre de paciente heterocigoto con la variante Hb D-Punjab
- PT : Amostra de sangue de doente heterozigótico com variante Hb D-Punjab
- SV : Blodprov från heterozygot patient med Hb D-Punjab variant
- GR : Δείγμα αίματος από ετερόζυγο ασθενή με παραλλαγή Hb D-Punjab
- HR : Uzorak krvi heterozigotnog bolesnika s varijantom Hb D-Punjab
- LT : Heterozigotinio paciento kraujo, kuriame yra Hb D-Punjab varianto, mėginys
- PL : Próba krwi od heterozygotycznego pacjenta z odmianą Hb D-Punjab
- RO : Probă de sânge de la pacient heterozigot cu varianta Hb D-Punjab
- CS : Uzorak krvi od heterozigotního pacienta sa Hb D-Punjab variantom
- HU : Heterozigóta beteg vérmintája Hb D-Punjab variánssal
- TR : Hb D-Punjab varyantı taşıyan heterozigot hastasına ait kan numunesi
- CZ : Vzorek krve heterozigotního pacienta s variantou Hb D-Punjab
- BG : Кръвна проба от хетерозиготен пациент с Hb D-Punjab вариант
- NO : Blodprobe fra heterozygot pasient med Hb D-Punjab variant
- DK : Blodprøve fra heterozygot patient med Hb D-Punjab-variant
- CN : 来自 Hb D-Punjab 变体杂合患者的血液样品
- RU : Образец крови от гетерозиготного пациента с вариацией Hb D-Punjab
- JP : Hb D-Punjab変異体を含むヘテロ接合体患者からの血液サンプル
- LV : Heterozigota pacienta asins paraugs ar Hb D-Punjab variantu
- SK : Vzorka krvi od heterozigotného pacienta s variantom Hb D-Punjab
- EE : Vereproov Hb D-Punjab variantiga heterosügotselt patsiendilt
- VN : Mẫu máu của bệnh nhân bị bệnh di hợp tử với biến thể Hb D-Punjab

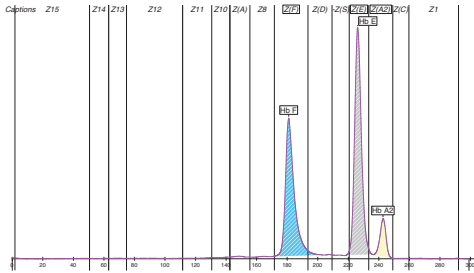
- Sang de patient hétérozygote avec variant delta Hb A2
- Blood sample from heterozygous patient with delta Hb A2 variant
- Blutprobe eines heterozygoten Patienten mit Delta-Hb A2-Variante
- Bloedmonster van heterozygote patiënt met delta Hb A2 variant
- Sangue di paziente eterozygote con variante delta Hb A2
- Sangre de paciente heterocigoto con variante delta Hb A2
- Amostra de sangue de doente heterozigótico com variante delta Hb A2
- Blodprov från heterozygot patient med delta Hb A2-variant
- Δείγμα αίματος από ετερόζυγο ασθενή με παραλλαγή δέλτα Hb A2
- Uzorak krvi heterozigotnog bolesnika s varijantom delta Hb A2
- Heterozigotinio paciento kraujo, kuriame yra delta Hb A2 varianto, mėginys
- Próba krwi od heterozygotycznego pacjenta z odmianą delta Hb A2
- Probă de sânge de la pacient heterozigot cu varianta delta Hb A2
- Uzorak krvi od heterozigotního pacienta sa delta Hb A2 varijantom
- Heterozigóta beteg vérmintája delta Hb A2 variánssal
- Delta Hb A2 varyantı taşıyan heterozigot hastasına ait kan numunesi
- Vzorek krve heterozigotního pacienta s variantou delta Hb A2
- Кръвна проба от хетерозиготен пациент с delta Hb A2 вариант
- Blodprobe fra heterozygot pasient med delta Hb A2 variant
- Blodprøve fra heterozygot patient med Hb A2-variant
- 来自 Hb A2 变体杂合患者的血液样品
- Образец крови от гетерозиготного пациента с вариацией delta Hb A2
- デルタHb A2変異体を含むヘテロ接合体患者からの血液サンプル
- Heterozigota pacienta asins paraugs ar delta Hb A2 variantu
- Vzorka krvi od heterozigotného pacienta s variantom Hb A2
- Vereproov delta Hb A2 variantiga heterosügotselt patsiendilt
- Mẫu máu của bệnh nhân bị bệnh di hợp tử với biến thể Hb A2 delta

SCHÉMAS / FIGURES

ABBILDUNGEN - FIGUREN - FIGURE - FIGURAS - BILDER - EIKONES - SLIKE - PAVEIKSLAI - RYSUNKI - FIGURI - ÁBRÁK - ŞEKİLLER - OBRÁZKY - ΦΙΓΥΡΗ - FIGURER - 插图 - РИСУНКИ - 図 - CIPARI - JOONISED - SƠ ĐỒ

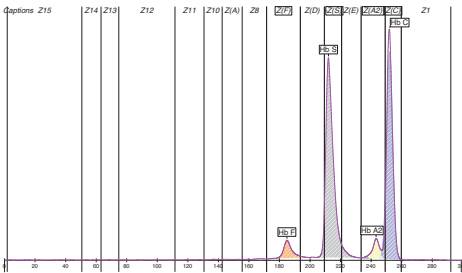
**CAP1 3 HEMOGLOBIN(E)
PROFILS ÉLECTROPHORÉTIQUES - ELECTROPHORETIC PATTERNS**

11



Sang de patient homozygote avec variant Hb E et fraction Hb F élevée
Blood sample from homozygous patient with Hb E variant and elevated Hb F

12



Sang de patient hétérozygote composite avec variants Hb S et Hb C
Blood sample from compound heterozygous patient with Hb S & Hb C variants

FR : Sang de patient homozygote avec variant Hb E et fraction Hb F élevée
GB : Blood sample from homozygous patient with Hb E variant and elevated Hb F
DE : Blutprobe eines homozygoten Patienten mit Hb E-Variante und erhöhtem Hb F
NL : Bloedmonster van homozygote patiënt met Hb E variant en verhoogd Hb F niveau
IT : Sangue di paziente omozigote con variante Hb E y la fracción Hb F elevada
ES : Sangre de paciente homocigoto con la variante Hb E y la fracción Hb F alta
PT : Amostra de sangue de doente heterocigótico com variante Hb E e Hb F elevada
SV : Blodprov från heterozygot patient med Hb E variant och förhöjt Hb F
GR : Δείγμα αίματος από ομόζυγο ασθενή με παραλλαγή Hb E και αυξημένο επίπεδο Hb F
HR : Uzorak krvi homocigotnog bolesnika s varijantom Hb E i povišenom vrijednošću Hb F
LT : Homocigotinio paciento kraujo, kuriame yra Hb E varianto ir padidėjęs Hb F kiekis, mėginys
PL : Próbk krwi od homocigotycznego pacjenta z odmianną Hb E i podwyższonym stężeniem Hb F
RO : Probă de sânge de la pacient heterocigot cu varianta Hb E și Hb F crescută
CS : Uzorak krvi od heterocigotnog pacijenta sa Hb E varijantom i povišenim Hb F
HU : Homozigóta beteg vérmintája Hb E variánsal és emelkedett Hb F-fel
TR : Hb E varyantı ve yüksek Hb F taşıyan homozigot hastasına ait kan numunesi
CZ : Vzorek kve homocigotního pacienta s variantou Hb E a zvýšením Hb F
BG : Кръвна проба от хомозиготен пациент с Hb E вариант и повишен Hb F
NO : Blodprobe fra homozygot pasient med Hb E variant og forhøyet Hb F
DK : Blodprobe fra homozygot patient med Hb E-variant og forhøjet Hb F
CN : 来自 Hb E 变体利 Hb F 升高余台患者的血液样品
RU : Образец крови от гомозиготного пациента с вариацией Hb E и повышенным уровнем Hb F
JP : Hb E変異体および高値のHb Fを含む本を接合患者からの血液サンプル
LV : Homozigota pacienta asiņš paraugs ar Hb E variantu un paaugstinātu Hb F līmeni
SK : Vzorka krvi od homozigotného pacienta s variantom Hb E a zvýšenou hladinou Hb F
EE : Vereproov Hb E varianti ja kõrgenenud Hb F-iga heterosigotseilt patsiendilt
VN : Máu mầu của bệnh nhân bị bệnh di hợp từ với biến thể Hb E và Hb F gia tăng

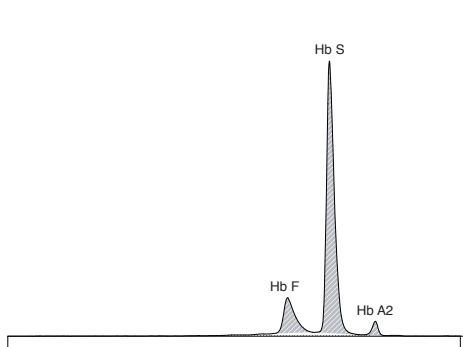
Sang de patient hétérozygote composite avec variants Hb S et Hb C
Blood sample from compound heterozygous patient with Hb S & Hb C variants
Blutprobe eines compound-heterozygoten Patienten mit Hb S- und Hb C-Varianten
Bloedmonster van samengestelde heterozygote patiënt met Hb S en Hb C varianten
Sangue di paziente eterozigote composto con varianti Hb S e Hb C
Sangre de paciente heterocigoto compuesto con las variantes Hb S e Hb C
Amostra de sangue de doente heterocigótico composto com variantes Hb S e Hb C
Blodprov från förenad heterozygot patient med Hb S & Hb C-varianter
Δείγμα αίματος από σύνθετο ετερόζυγο ασθενή με παραλλαγές Hb S & Hb C
Uzorak krvi složenog heterocigotnog bolesnika s varijantama Hb S i Hb C
Paciento kraujo, kuriame yra heterocigotinių junginių ir Hb S bei Hb C variantų, mėginys
Próbka krwi od heterocigotycznego pacjenta z jednoczesną obecnością odmiann Hb S oraz Hb C
Probă de sânge de la pacient heterocigot compus cu variantele Hb S și Hb C
Uzorak krvi od složenog heterocigotnog pacijenta sa Hb S & Hb C varijantama
Összetett heterozigóta beteg vérmintája Hb S és Hb C variánsokkal
Hb S ve Hb C varyantlarını taşıyan bileşik heterozigot hastasına ait kan numunesi
Vzorek krvi sdrúženého heterocigotního pacienta s variantami Hb S a Hb C
Кръвна проба от пациент със съставна хетерозиготност с Hb S и Hb C варианти
Blodprobe fra sammensatt heterozygot pasient med Hb S og Hb C-varianter
Blodprobe fra heterozygot patient med Hb S- og Hb C-varianter
来自 Hb S & Hb C 变体余台患者的血液样品
Образец крови от компунд-heterozigota с вариациями Hb S и Hb C
Hb SおよびHb C変異体を含む複合ヘテロ接合患者からの血液サンプル
Kompaundna heterozigota pacienta asiņš paraugs ar Hb S un Hb C variantu
Vzorka krvi od heterozigotného pacienta s variantami Hb S a Hb C
Vereproov Hb S ja Hb C variantidega ühend-heterosigotseilt patsiendilt
Máu mầu của bệnh nhân bị bệnh di hợp từ kép với biến thể Hb S & Hb C

SCHÉMAS / FIGURES

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CAP1 3 HEMOGLOBIN(E)
 PROFILS ÉLECTROPHORÉTIQUES - ELECTROPHORETIC PATTERNS

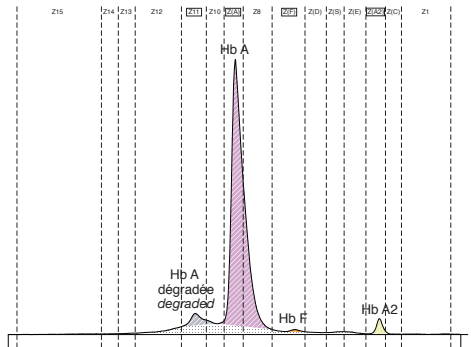
13



Sang de patient homozygote avec Hb F et variant Hb S
 Blood sample from homozygous patient with Hb F and Hb S variant

- FR : Sang de patient homozygote avec Hb F et variant Hb S
- GB : Blood sample from homozygous patient with Hb F and Hb S variant
- DE : Blutprobe eines homozygoten Patienten mit Hb F- und Hb S-Variante
- NL : Bloedmonster van homozygote patiënt met Hb F en Hb S-variant
- IT : Sangue di paziente omozigote con Hb F e variante Hb S
- ES : Sangre de paciente homocigoto con Hb F y la variante Hb S
- PT : Amostra de sangue de doente homocigótico com variantes Hb F e Hb S
- SV : Blodprov från homozygot patient med Hb F- och Hb S-variant
- GR : Δείγμα αίματος από ομοζυγό ασθενή με παραλλαγή Hb F και Hb S
- HR : Uzorak krvi homozigotnog bolesnika s varijantama Hb F i Hb S
- LT : Homozigotinio paciento kraujo, kuriame yra Hb F bei Hb S variantų, mėginys
- PL : Próba krwi od homozigotnego pacjenta z odmiąną Hb F i Hb S.
- RO : Probă de sânge de la pacient heterozigot cu variantele Hb F și Hb S
- CS : Uzorak krvi od heterozigotnog pacijenta sa Hb F i Hb S varijantom
- HU : Homozigóta beteg vérmintája Hb F-fei és Hb S variánsal
- TR : Hb F ve Hb S varyantlarını taşıyan homozigot hastasına ait kan numunesi
- CZ : Vzorek krve homozigotního pacienta s variantami Hb F a Hb S
- BG : Кръвна проба от хомозиготен пациент с Hb F и Hb S вариант
- NO : Blodprobe fra homozygot pasient med Hb F og Hb S variant
- DK : Blodprobe fra heterozygot patient med Hb F- og Hb S-variant
- CN : 来自 Hb F 和 Hb S 变体杂合患者的血液样品
- RU : Образец крови от гомозиготного пациента с вариациями Hb F и Hb S
- JP : Hb FおよびHb S変異体を含むホモ接合患者からの血液サンプル
- LV : Heterozigota pacienta asins paraugs ar Hb F un Hb S variantu
- SK : Vzorka krvi od homozygotného pacienta s variantmi Hb F a Hb S
- EE : Vereproov Hb F ja Hb S variantiga homosiigotseilt patsiidilt
- VN : Mẫu máu của bệnh nhân bị bệnh di hợp tử với biến Hb thF và Hb S

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Sang avec Hb A dégradée (Hb A3) et Hb F faible
 Blood sample with degraded Hb A (Hb A3) and faint Hb F

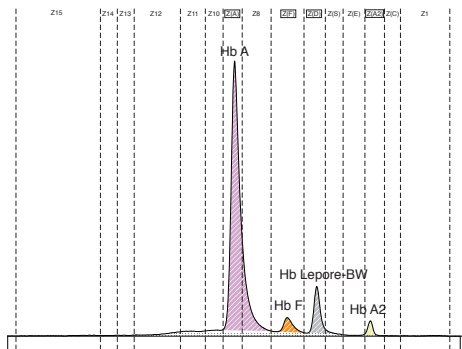
- Sang avec Hb A dégradée (Hb A3) et Hb F faible
- Blood sample with degraded Hb A (Hb A3) and faint Hb F
- Blutprobe mit degradiertem Hb A (Hb A3) und schwachem Hb F
- Bloedmonster met afgebroken Hb A (Hb A3) en nauwelijks waarneembare Hb F
- Sangue con Hb A degradata (Hb A3) e Hb F bassa
- Sangre con Hb A degradada (Hb A3) y Hb F débil
- Amostra de sangue com Hb A (Hb A3) degradada e Hb F baixa
- Blodprov med nedbrutet Hb A (Hb A3) och svagt Hb F
- Αδείγμα αίματος με αποσυντεθειμένη Hb A (Hb A3) και αμυδρή Hb F
- Uzorak krvi s degradiranim Hb A (Hb A3) i slabim Hb F
- Kraujo, kuriame yra suskaidusio Hb A (Hb A3) ir silbuko Hb F mėginys
- Próbka krwi z rozłożoną Hb A (Hb A3) i śladową obecnością Hb F
- Probă de sânge cu Hb A degradată (Hb A3) și Hb F slabă
- Uzorak krvi sa degradiranim Hb A (Hb A3) i niskim Hb F
- Vérminta degradálódott Hb A-val (Hb A3) és halvány Hb F-fei
- Indirgenmiş Hb A (Hb A3) ve belirsiz/zayıf Hb F içeren kan numunesi
- Vzorek krve s degradovaným Hb A (Hb A3) a slabým Hb F
- Кръвна проба с разграден Hb A (Hb A3) и малко количество Hb F
- Blodprobe med degradert Hb A (Hb A3) og svakt Hb F
- Blodprobe med nedbrudt Hb A (Hb A3) og svag Hb F
- Hb A (Hb A3) 降低和Hb F虚弱的血液样品
- Образец крови с подвергнувшимся разложению Hb A (Hb A3) и низким Hb F
- 劣化したHb A (Hb A3) および僅少なHb Fを含む血液サンプル
- Asins paraugs ar noārdētu Hb A (Hb A3) un nelielu Hb F daudzumu
- Vzorka krvi s degradovaným Hb A (Hb A3) a nevýrazným Hb F
- Vereproov lagunenud Hb A (Hb A3) ja nõrga Hb F-ga
- Mẫu máu có Hb A (Hb A3) suy giảm và Hb F yếu

SCHEMAS / FIGURES

ABBILDUNGEN - FIGUREN - FIGURE - FIGURAS - BILDER - EIKONES - SLIKE - PAVEIKSLAI - RYSUNKI - FIGURI -
 ÁBRÁK - ŞEKİLLER - OBRÁZKY - ΦΙΓΥΡΗ - FIGURER - 插图 - РИСУНКИ - 図 - CIPARI - JOONISED - SƠ ĐỒ

CAPI 3 HEMOGLOBIN(E)
 PROFILS ÉLECTROPHORÉTIQUES - ELECTROPHORETIC PATTERNS

15



Sang de patient hétérozygote avec variant Hb Lepore-Boston-Washington
 Blood sample from heterozygous patient with Hb Lepore-Boston-Washington variant

FR : Sang de patient hétérozygote avec variant Hb Lepore-Boston-Washington
 GB : Blood sample from heterozygous patient with Hb Lepore-Boston-Washington variant
 DE : Blutprobe eines heterozygoten Patienten mit Hb Lepore-Boston-Washington-Variante
 NL : Bloedmonster van heterozygote patiënt met Hb Lepore-Boston-Washington variant
 IT : Sangue di paziente eterozigote con variante Hb Lepore-Boston-Washington
 ES : Sangre de paciente heterocigoto con la variante Hb Lepore-Boston-Washington
 PT : Amostra de sangue de doente heterozigótico com variante Hb Lepore-Boston-Washington
 SV : Blodprov från heterozygot patient med Hb Lepore-Boston-Washington-variant
 GR : Δείγμα αίματος από ετερόζυγο ασθενή με παραλλαγή Hb Lepore-Boston-Washington
 HR : Uzorak krvi heterozigotnog bolesnika s varijantom Hb Lepore-Boston-Washington
 LT : Heterozigotinio paciento kraujo, kuriame yra Hb Lepore-Boston-Washington varianto, kraujo mėginys
 PL : Próba krwi od heterozygotycznego pacjenta z odmianą Hb Lepore-Boston-Washington
 RO : Probă de sângue de la pacient heterozigot cu varianta Hb Lepore-Boston-Washington
 CS : Uzorak krvi od heterozigotnog pacijenta sa Hb Lepore-Boston-Washington varijantom
 HU : Heterozigóta beteg vérmintája Hb Lepore-Boston-Washington variánsa
 TR : Hb Lepore-Boston-Washington varyantı taşıyan heterozigot hastasına ait kan numunesi
 CZ : Vzorek krve heterozigotního pacienta s variantou Hb Lepore-Boston-Washington
 BG : Кръвна проба от хетерозиготен пациент с Hb Lepore-Boston-Washington вариант
 NO : Blodprøve fra heterozygot pasient med Hb Lepore-Boston-Washington variant
 DK : Blodprøve fra heterozygot patient med Hb Lepore-Boston-Washington-variant
 CN : 来自 Hb Lepore-Boston-Washington 变异杂合患者的血液样品
 RU : Образец крови от гетерозиготного пациента с вариацией Hb Lepore-Boston-Washington
 JP : Hb Lepore-Boston-Washington変異体を含むヘテロ接合体からの血液サンプル
 LV : Heterozigota pacienta asins paraugs ar Hb Lepore-Boston-Washington variantu
 SK : Vzorka krvi od heterozigotného pacienta s variantom Hb Lepore-Boston-Washington
 EE : Vereproov Hb Lepore-Boston-Washingtoni variantiga heterosügootselt isegiendil
 VN : Mẫu máu của bệnh nhân bị bệnh di truyền từ với biến thể Hb Lepore-Boston-Washington

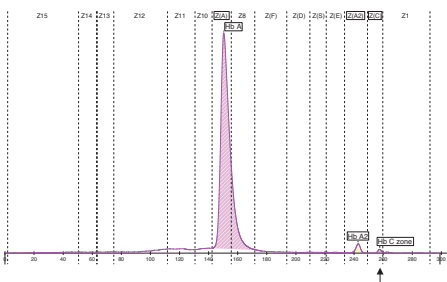
SCHÉMAS / FIGURES

ABBILDUNGEN - FIGUREN - FIGURE - FIGURAS - BILDER - ΕΙΚΟΝΕΣ - SLIKE - PAVEIKSLAI - RYSUNKI - FIGURI - ÁBRÁK - ŞEKİLLER - OBRÁZKY - ΦΙΓΥΡΗ - FIGURER - 插图 - РИСУНКИ - 図 - CIPARI - JOONISED - 図

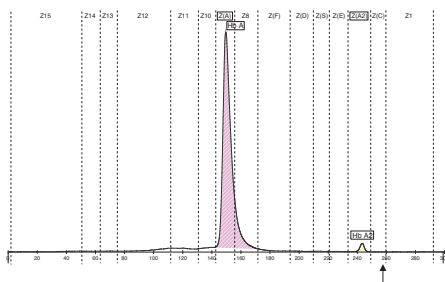
CAP1 3 HEMOGLOBIN(E)
 PROFILS ÉLECTROPHORÉTIQUES - ELECTROPHORETIC PATTERNS

16

Fraction supplémentaire en zone de migration Z(C) (protéines plasmatiques)
 Additional fraction in Z(C) migration zone (plasmatic proteins)



Analyse du sang total
 Whole blood analysis



Analyse des globules rouges correspondants
 Analysis of corresponding red blood cells

FR : Fraction supplémentaire en zone de migration Z(C) (protéines plasmatiques)

GB : Additional fraction in Z(C) migration zone (plasmatic proteins)

DE : Zusätzliche Fraktion in der Z(C)-Migrationszone (plasmatische Proteine)

NL : Bijkomende fractie in Z(C) migratiezone (plasma-eiwitten)

IT : Frazione addizionale in zona di migrazione Z(C) (proteine plasmatiche)

ES : Fracción adicional en la zona de migración Z(C) (proteínas plasmáticas)

PT : Fração adicional na zona de migração Z(C) (proteínas plasmáticas)

SV : Extra fraktion i Z(C) migreringszon (plasmatiska proteiner)

GR : Πρόσθετο κλάσμα σε ζώνη μετακίνησης Z(C) (πρωτεΐνες πλάσματος)

HR : Dodatna frakcija u zoni migracije Z(C) (plazmatski proteini)

LT : Papildoma frakcija Z(C) migravimo zonoje (plazmatiniai baltymai)

PL : Dodatkowa frakcja w strefie migracji Z(C) (białka plazmatyczne)

RO : Frație suplimentară în zona de migrare Z(C) (proteine plasmactice)

CS : Dodatná frakcija u zoni migracije Z(C) (plazmatski proteini)

HU : További frakció a Z(C) migrációs zónában (plazmafehérjék)

TR : Z(C) migrasyon bölgesinde ek fraksiyon (plazma proteinleri)

CZ : Další frakce v migrační zóně Z(C) (plazmatické proteiny)

BG : Допълнителна фракция в зона на миграция Z(C) (плазмени протеини)

NO : Tilleggsfraksjon i Z(C) migrasjonszone (plasmatiske proteiner)

DK : Ekstra fraktion i Z(C) migrationszonen (plasma-proteiner)

CN : Z(C) 电泳区的其他底带 (血浆蛋白)

RU : Дополнительная фракция в зоне миграции Z(C) (плазматические белки)

JP : Z(C)泳動領域における追加フラクション (血漿タンパク)

LV : Papildu sadaļa Z(C) migrācijas zonā (plazmas olbaltumvielas)

SK : Dodatočná frakcia v zóne migrácie Z(C) (plazmatické proteíny)

EE : Lisafraktsioon Z(C) migratsioonisoonis (plasmavalgud)

VN : Phần đoạn bổ sung trong vùng di chuyển Z(C) (protein huyết tương)

Analyse du sang total

Whole blood analysis

Vollblutanalyse

Voilbloedanalyse

Analisi su sangue intero

Analisis de sangre total

Analise do sangue total

Hei blodanalyse

Ανάλυση ολικού αίματος

Analiza pune krvi

Visos sudėties kraujo analizė

Analiza krvi pehne

Hemoleucograma completă

Analiza cele krvi

Teljesvér-vizsgálat

Tam kan analizi

Analiza pline krve

Analiz na čla krvi

Fullstendig blodanalyse

Fülldblodanalyse

全血分析

Analiz celynoy krovi

全血分析

Plina asins aina

Analiza pneh krvi

Täisvere analüüs

Phân tích máu toàn phần

Analyse des globules rouges correspondants

Analysis of corresponding red blood cells

Analise der entsprecheden roten Blutkörperchen

Analyse van overeenkomstige rode bloedlichaampjes

Analisi su globuli rossi corrispondenti

Analisis de los glóbulos rojos correspondientes

Analise dos eritrócitos correspondentes

Analys av motsvarande röda blodkroppar

Ανάλυση των αντίστοιχων ερυθροκυττάρων

Analiza odgovarajućih crvenih krvnih stanica

Atitinkami eritrocitų analizė

Analiza odpowiednich czerwonych czerwonyc

Analiza eritrocitelor aferente

Analiza odgovarajućih crvenih krvnih ćelija

Megfelelő vörösvértestek vizsgálata

Ilgili kırmızı kan hücrelerinin analizi

Analiza odgovarajucih červenych krvinek

Analiz na съответните червени кръвни телца

Analise av tilsvarende røde blodlegemer

Analise af tilsvarende røde blodceller

分析相应的红细胞

Analiz sootvetstvayushchey eritrotsitoev

分析相应的红细胞

Atbilstošo sarkano asins ķermeņu šūnu analīze

Analiza zodpovedajucich červenych krvinek

Vastavate punaste vereleible analüüs

Phân tích tế bào hồng cầu tương ứng

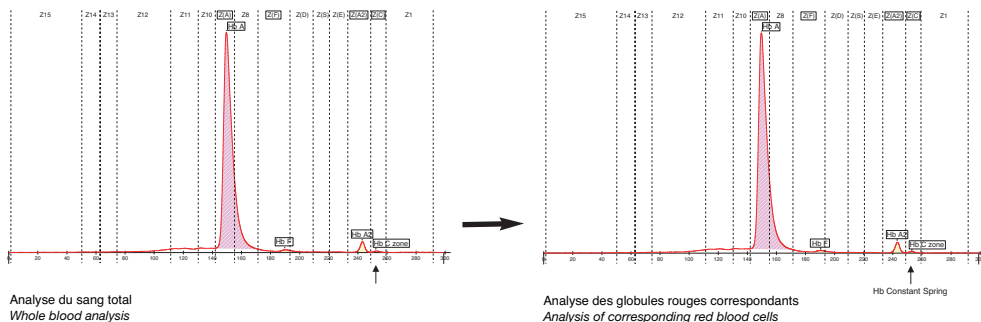
SCHÉMAS / FIGURES

ABBILDUNGEN - FIGUREN - FIGURE - FIGURAS - BILDER - ΕΙΚΟΝΕΣ - SLIKE - PAVEIKSLAI - RYSUNKI - FIGURI -
 ÁBRÁK - ŞEKİLLER - OBRÁZKY - ΦΙΓΥΡΗ - FIGURER - 插图 - РИСУНКИ - 図 - CIPARI - JOONISED - SƠ ĐỒ

CAPI 3 HEMOGLOBIN(E)
 PROFILS ÉLECTROPHORÉTIQUES - ELECTROPHORETIC PATTERNS

17

Fraction supplémentaire en zone de migration Z(C) (Hb Constant spring)
 Additional fraction in Z(C) migration zone (Hb Constant spring)



FR : Fraction supplémentaire en zone de migration Z(C) (Hb Constant spring)
 GB : Additional fraction in Z(C) migration zone (Hb Constant spring)
 DE : Zusätzliche Fraktion in der Z(C)-Migrationszone (Hb Constant Spring)
 NL : Bijkomende fractie in Z(C) migratiezone (Hb Constant spring)
 IT : Frazione addizionale in zona di migrazione Z(C) (Hb Constant spring)
 ES : Fracción adicional en la zona de migración Z(C) (Hb Constant spring)
 PT : Fração adicional na zona de migração Z(C) (Hb Constant spring)
 SV : Extra fraktion i Z(C) migreringszon (Hb Constant spring)
 GR : Πρόσθετο κλάσμα σε ζώνη μετακίνησης Z(C) (Hb Constant spring)
 HR : Dodatna frakcija u zoni migracije Z(C) (Hb Constant Spring)
 LT : Papildoma frakcija Z(C) migravimo zonoje (Hb Constant Springas)
 PL : Dodatkowa frakcja w strefie migracji Z(C) (Hb Constant spring)
 RO : Frație suplimentară în zona de migrare Z(C) (Hb Constant Spring)
 CS : Dodatná frakcija u zoni migracije Z(C) (Hb Constant spring)
 HU : További frakció a Z(C) migrációs zónában (Hb Constant Spring)
 TR : Z(C) migrasyon bölgesinde ek fraksiyon (Hb Sabit Yay)
 CZ : Další frakce v migrační zóně Z(C) (Hb Constant spring)
 BG : Допълнителна фракция в зона на миграция Z(C) (Hb с удължена еверига)
 NO : Tilleggsfraksjon i Z(C) migrasjonszone (Hb konstant fjær)
 DK : Ekstra fraktion i Z(C)-migrationszonen (Hb-konstant spring)
 CN : Z(C) 电泳区的其他区带 (Hb Constant spring)
 RU : Дополнительная фракция в зоне миграции Z(C) (Гемоглобин Констант-Спринг)
 JP : Z(C)泳動領域における追加ラクション (Hb Constant spring)
 LV : Papildu sadaja Z(C) migrācijas zonā (Hb atsauces konstante)
 SK : Dodatočná frakcia v zóne migrácie Z(C) (Hb Constant spring)
 EE : Lisafraktsioon Z(C) migratsioonitoonis (Hb Constant spring)
 VN : Phần đoạn bổ sung trong vùng di chuyển Z(C) (Hb Constant spring)

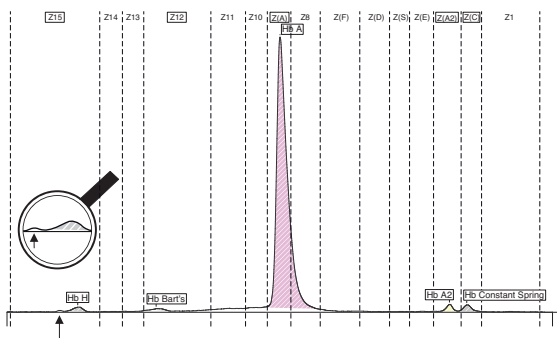
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CAPI 3 HEMOGLOBIN(E)
 PROFILS ÉLECTROPHORÉTIQUES - ELECTROPHORETIC PATTERNS

18

Fraction supplémentaire en zone de migration Z15
 Additional fraction in Z15 migration zone



FR : Fraction supplémentaire en zone de migration Z15

GB : Additional fraction in Z15 migration zone

DE : Zusätzliche Fraktion in der Z15-Migrationszone

NL : Bijkomende fractie in Z15 migratiezone

IT : Frazione addizionale in zona di migrazione Z15

ES : Fracción adicional en la zona de migración Z15

PT : Fração adicional na zona de migração Z15

SV : Extra fraktion i Z15 migreringszon

GR : Πρόσθετο κλάσμα σε ζώνη μετακίνησης Z15

HR : Dodatna frakcija u zoni migracije Z15

LT : Papildoma frakcija Z15 migravimo zonoje

PL : Dodatkowa frakcja w strefie migracji Z15

RO : Fracție suplimentară în zona de migrare Z15

CS : Dodatná frakcija u zoni migrácie Z15

HU : További frakció a Z15 migrációs zónában

TR : Z15 migrasyon bölgesinde ek fraksiyon

CZ : Další frakce v migrační zóně Z15

BG : Допълнителна фракция в зона на миграция Z15

NO : Tilleggsfraksjon i Z15 migrasjonszone

DK : Ekstra fraktion i Z15-migrationszonen

CN : Z15 电泳区的其他区带

RU : Дополнительная фракция в зоне миграции Z15

JP : Z15泳動領域における追加フラクション

LV : Papildu sadāja Z15 migrācijas zonā

SK : Dodatočná frakcia v zóne migrácie Z15

EE : Lisafraktsioon Z15 migratsioonitsoonis

VN : Phần đoạn bổ sung trong vùng di chuyển Z15



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