



View Protocol

ChromoQuant® QF PCR kit **Optima PLUS**

For detection of trisomy in chromosomes 15, 16 and 22

Proven technology QF-PCR

- Fast analysis Turnaround reporting time is less than 24
- **High Throughput PCR** based system. Automatable for cost efficient analysis
- High specificity 22 + 15 markers
- **Combination with ChromoQuant Optima STaR:** ChromoQuant STaR **PLUS**
- GeneMapper and GeneMarker panels

Optima STaR PLUS

For detection of aneuploidy in chromosomes 13,15,16,18,21,22 and X/Y

The ChromoQuant Optima PLUS QF-PCR kit is used for fast and accurate diagnosis of the most common aneuploidies in miscarriages

Key advantages

- High number of genetic markers for maximal speed and accuracy
- **QF-PCR Technology**
- GeneMapper and GeneMarker panel templates are available
- No tissue culture
- Detection of maternal contamination eliminates risk of misdiagnosis
- Flexibel solution in combination with other ChromoQuant Optima kits.

CE marked IVD kit

High flexibility

ChromoQuant® is CE marked in accordance with the Directive 98/79/EC. ChromoQuant® is produced under quality certificate ISO 13485.

ChromoQuant Optima PLUS with markers for

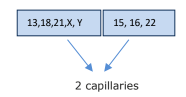
separately or in combination with ChromoQuant STaR Optima or other

ChromoQuant Optima kits as preferred.

chromosomes 15, 16 and 22 can be purchased

High specificity

ChromoQuant Optima PLUS, 15 markers in total for Chr. 15, 16 and 22, single tube ChromoQuant Optima STaR PLUS, 37 markers in total for Chr. 13, 15, 16, 18, 21, 22, X and Y, duplex test.





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CyberGene AB

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PCR based products

is active in the MedTech field

by developing, manufacturing and selling In Vitro diagnostic



Intended Use Optima PLUS ChromoQuant Optima PLUS: For detection of trisomy

in chromosomes 15, 16 and 22. Used for analysing

reasons for miscarriages.

Combi kit Optima STaR

PLUS

ChromoQuant Optima STaR PLUS: For detection of aneuploidy in chromosomes 13,15,16,18,21,22 and X/Y

Ready to use Add only DNA. Ready for PCR.

Optima PLUS QF-PCR test for diagnosing aneuploidy in

Chr. 15, 16 and 22. 15 markers in total.

Optima STaR PLUS QF-PCR test for diagnosing aneuploidy in

Chr. 13, 15, 16, 18, 21, 22, XY.

37 markers in total.

PCR:

Optima PLUS The markers for chromosomes 15, 16 and 22 are

amplified in one tube.

Optima STaR PLUS The markers for Chr. 13, 18, 21, XY and Chr. 15, 16,

22 are amplified separated in two tubes.

Number of markers:

Markers Optima PLUS 5 STR markers for each Chr. 15, 16 and 22

Markers Optima STaR PLUS 5 markers for Chr. 13 and 18

6 markers for Chr. 21

6 markers for X and Y. Marker for Turner X0 included. 5 STR markers for each chromosome 15, 16 and 22

Kit size:

Optima PLUS 26 tests (Optima PLUS, P/N 531.001-26)

Optima STaR PLUS 26+26 (Optima STaR PLUS, P/N 514.531-26)

CE-marked for IVD use Yes

Detection format Capillary Electrophoresis

CE-marked for IVD use Yes

Validated Sequencers ABI 310, 3100, 3130, 3730, 3500

Data Interpretation GeneMapper and GeneMarker plugins are available

Detection format Capillary Electrophoresis

Complies with Best Yes

Practice Guidelines

STANTEM CEATING ISO 13485

ChromoQuant® has been thoroughly validated in hospital clinics. ChromoQuant® was clinically introduced in early 2004 and is used world wide.

India Contact: Life Technologies (India) Pvt. Ltd.