

sFlt-1  
PIGF



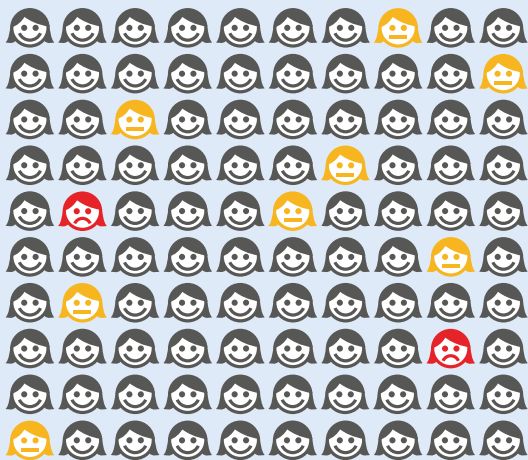
## Pre-eclampsia management with biomarkers

Improved pre-eclampsia diagnosis and prognosis of adverse outcome with PIGF and sFlt-1 after week 20 of gestation

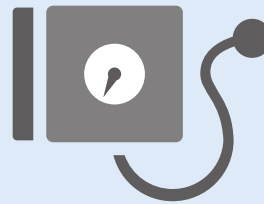
# Biomarkers for pre-eclampsia management

## Improving the diagnostic tools for pre-eclampsia evaluation

Pre-eclampsia is a progressive, pregnancy-related disorder with severe complications for mother and child. A timely diagnosis is needed in order to prevent maternal and fetal morbidity or mortality. In the absence of a specific therapy other than delivery the main objective of frequent patient monitoring is to detect deterioration of a patient's condition and to counteract maternal and fetal risk.



Diagnostic standard for pre-eclampsia ...



**Blood pressure**



**Proteinuria**



**10%** of pregnant women show unspecific signs and symptoms of pre-eclampsia



Only **one fifth of them** are actually developing pre-eclampsia<sup>1</sup>

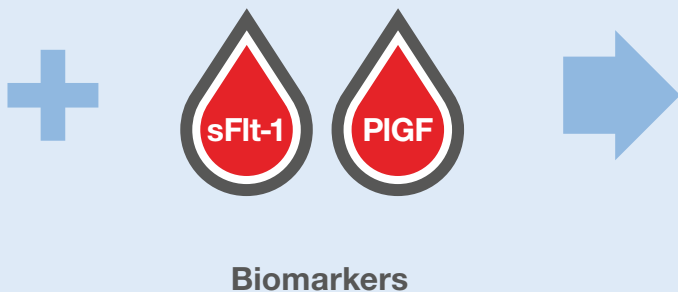
The “gold standard” for pre-eclampsia diagnosis – assessment of blood pressure and proteinuria – offers only a **poor sensitivity and specificity** with regards to origin of disease and prediction of maternal and perinatal outcome.<sup>2,3</sup>

## Serum sFlt-1 and PIGF determination adds significant clinical benefit to standard procedures

Determination of the biomarkers sFlt-1 (soluble FMS-like Tyrosine Kinase) and PIGF (Placental Growth Factor) in maternal blood have shown to significantly improve risk stratification among women presenting for pre-eclampsia evaluation.

The high sensitivity assays Thermo Scientific™ B·R·A·H·M·S™ sFlt-1 KRYPTOR™ and Thermo Scientific B·R·A·H·M·S PIGF plus KRYPTOR detect serum levels of both biomarkers reliably throughout pregnancy and thus improve pre-eclampsia management.

... significantly improved by sFlt-1 and PIGF serum measurement



Clearly **differentiating** between pre-eclampsia and other forms of hypertensive disorders<sup>4,5</sup>



Reliable **prognosis** of adverse outcome in women with suspected pre-eclampsia<sup>6,6</sup>



Offering **potential savings** in hospital costs and resource use<sup>7</sup>

**Measuring sFlt-1 and PIGF starting in mid pregnancy in women with suspected pre-eclampsia significantly improves the current evaluation of patients – for a better patient management and improved care.**

# Pre-eclampsia diagnosis and prognosis of adverse outcome

## The added value of sFlt-1 and PIGF

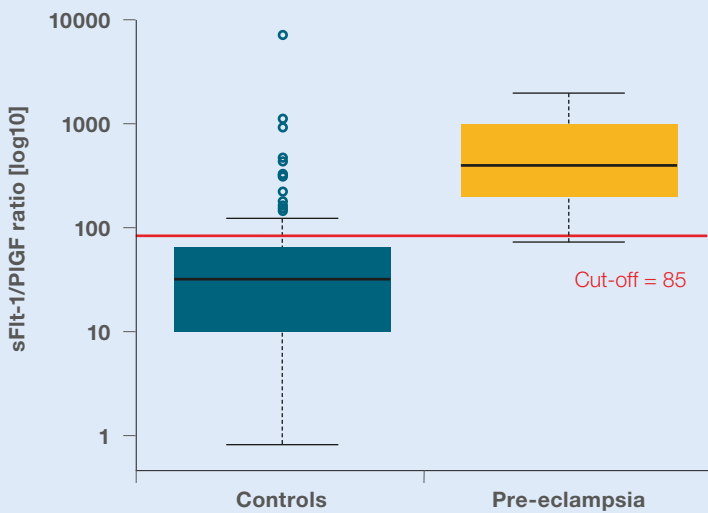
### Improved diagnosis of pre-eclampsia with sFlt-1/PIGF ratio

Many studies have proved the additional benefit of the sFlt-1/PIGF ratio in diagnosing pre-eclampsia:

- In women presenting with hypertension, the sFlt-1/PIGF ratio is able to **distinguish between those who will develop pre-eclampsia and those with chronic or gestational hypertension**. Women with pre-eclampsia have a significantly higher sFlt-1/PIGF ratio than women with other hypertensive disorders or controls.<sup>4,5</sup>
- The addition of sFlt-1/PIGF ratio to Doppler ultrasound measurement **improves the sensitivity and specificity** in diagnosing pre-eclampsia compared to the Doppler measurement alone.<sup>6</sup>

- Measurement of sFlt-1 and PIGF levels in maternal serum, starting in mid pregnancy, can **confirm pre-eclampsia diagnosis**, with the sFlt-1/PIGF ratio having a superior diagnostic ability compared to either of the biomarkers alone.<sup>8,9</sup>

**Therefore, the sFlt-1/PIGF ratio is a valuable additional tool for confirming or excluding the diagnosis of pre-eclampsia.**



PIGF and sFlt-1 were measured on KRYPTOR in parallel on samples from pregnant women with normal pregnancy outcome and patients with pre-eclampsia. At a cut-off of 85 for the sFlt-1/PIGF ratio, the sensitivity was calculated at 95% and the specificity at 84% for diagnosing pre-eclampsia. Latest studies show identical clinical performance and high accuracy in diagnosing pre-eclampsia when applying recently published cut-offs by using the sFlt-1/PIGF ratio on KRYPTOR.<sup>6,10,11</sup>

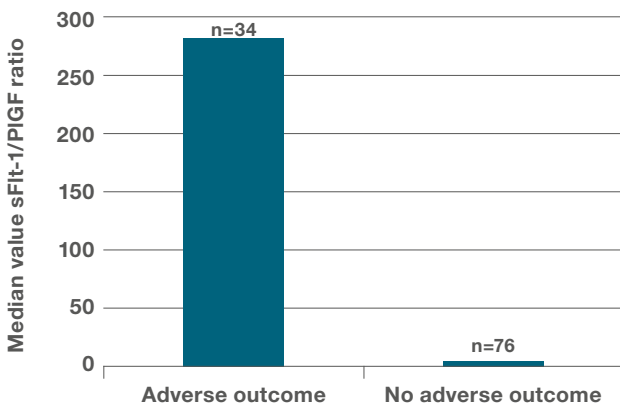
**The higher the sensitivity of a test the more women with pre-eclampsia are identified correctly and can be advised for closer monitoring.**

**Figure 1** Improved pre-eclampsia diagnosis with sFlt-1/PIGF ratio<sup>12</sup>



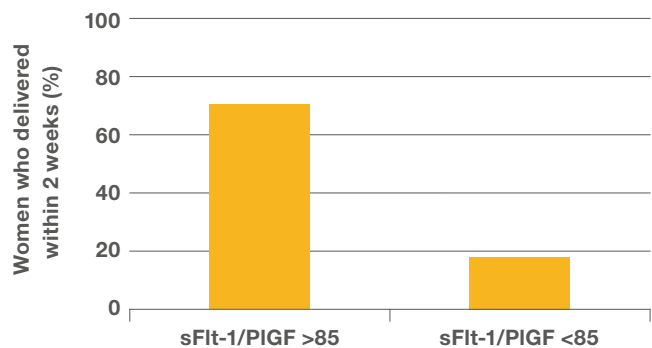
### Prognosis of adverse outcome with sFit-1/PIGF ratio

Recent studies showed that **women with any subsequent adverse outcome** in addition to hypertension had a significantly higher sFit-1/PIGF ratio than those women without, especially when presenting before week 34 (Figure 2).<sup>5,6</sup>



**Figure 2** Prediction of adverse outcome with sFit-1/PIGF ratio in women presenting < 34 weeks' gestation<sup>6</sup>

**Women who needed to be delivered within the next 2 weeks** after presentation had a significantly higher sFit-1/PIGF ratio than women who could continue with their pregnancy (Figure 3).<sup>5,6</sup>



**Figure 3** Prediction of duration of pregnancy with sFit-1/PIGF ratio in women presenting < 34 weeks' gestation<sup>6</sup>

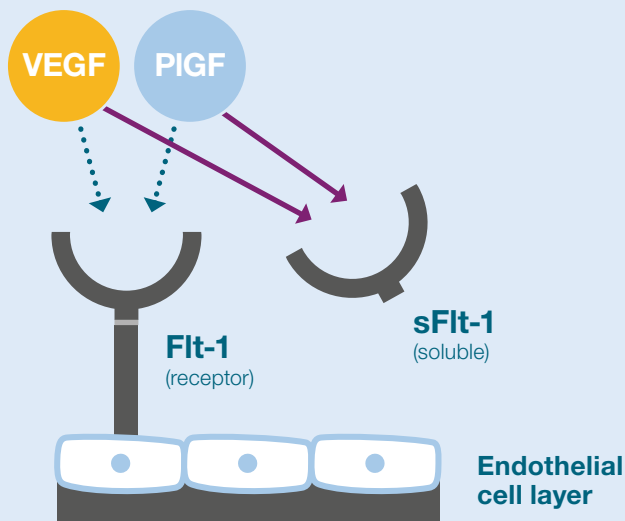
**The sFit-1/PIGF ratio is also a potent predictor for subsequent maternal and fetal adverse outcome in women already diagnosed with pre-eclampsia and can support clinical decisions.**

# The role of angiogenic factors

Biomarker levels correlate with severity of disease

## sFlt-1 and PlGF are counterparts

Although the cause of pre-eclampsia remains unclear, it is likely that the syndrome is initiated by an imbalance of angiogenic factors secreted by the placenta that induce endothelial dysfunction.



**sFlt-1** is a truncated form of the VEGF receptor Flt-1, circulating freely in the blood. sFlt-1 is produced in the placenta and secreted into the bloodstream, where it binds VEGF and PlGF with high affinity and therefore neutralizes their effects.<sup>8</sup>

**PlGF** belongs to the Vascular Endothelial Growth Factors (VEGF) family, promoting proliferation and survival of endothelial cells and inducing vascular permeability.<sup>13</sup>

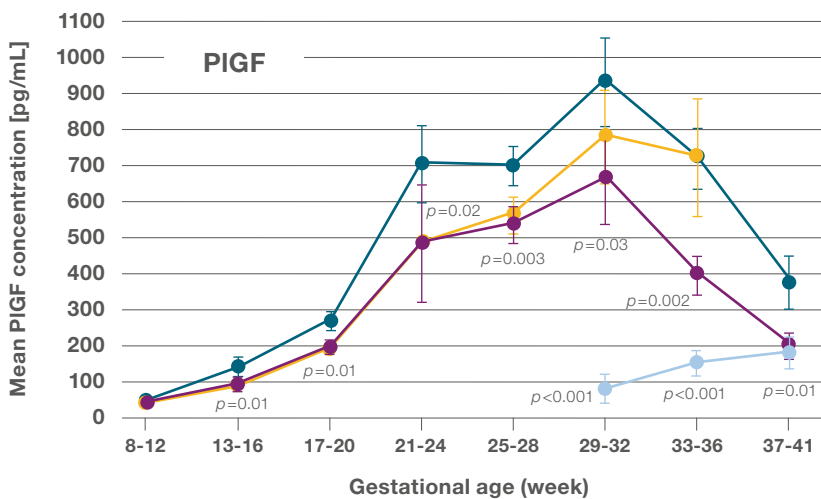
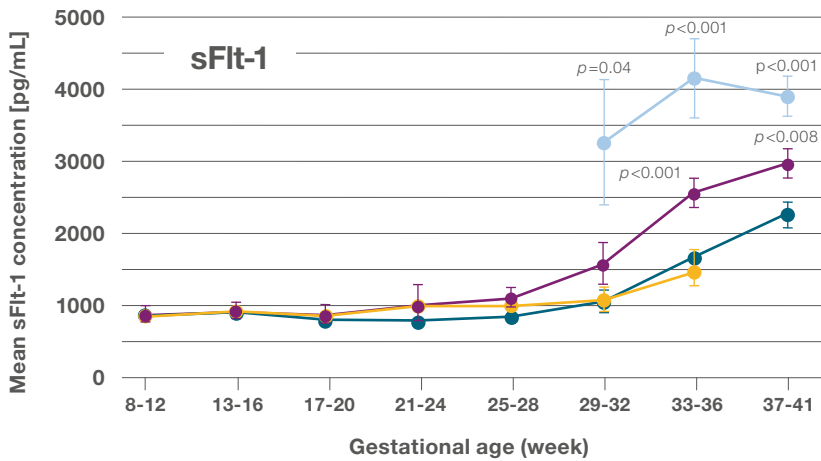
••▶ Signal transduction (healthy)

▶ Signal transduction inhibited

**Figure 4** sFlt-1 acts as potent antagonist of PlGF and VEGF by adhering to the receptor-binding domains, thus preventing interaction with endothelial receptors and inducing endothelial dysfunction







## Angiogenic factors during pregnancy

### Normal pregnancy

During pregnancy, sFlt-1 levels are stable until weeks 20-24, when they rise steadily until delivery. In contrast, PIGF levels increase progressively in first and second trimester and decrease towards term.<sup>13</sup>

### Pre-eclamptic pregnancy

In women with pre-eclampsia, sFlt-1 levels are significantly increased while concentrations of circulating free PIGF are significantly decreased.<sup>13,14</sup> In contrast to PIGF where the difference between healthy and pre-eclamptic pregnancies is measurable throughout pregnancy, sFlt-1 levels only start to separate after week 20.

- Controls
- Women who had pre-eclampsia >5 weeks later
- Women who later had pre-eclampsia
- Women with clinical pre-eclampsia

**Figure 5** Mean sFlt-1 and PIGF concentrations of healthy women and those women who later developed pre-eclampsia<sup>13</sup>

**Measuring maternal serum concentrations of sFlt-1 and PIGF can differentiate healthy women from women with pre-eclampsia.<sup>9,15</sup> Changes in sFlt-1 and PIGF levels also reflect the severity of the disease: early-onset pre-eclampsia is associated with greater changes compared to late-onset pre-eclampsia.<sup>16</sup>**

# Pre-eclampsia management throughout pregnancy

## Improving the outcomes for mother and child

### PIGF and PAPP-A: First trimester screening for timely intervention

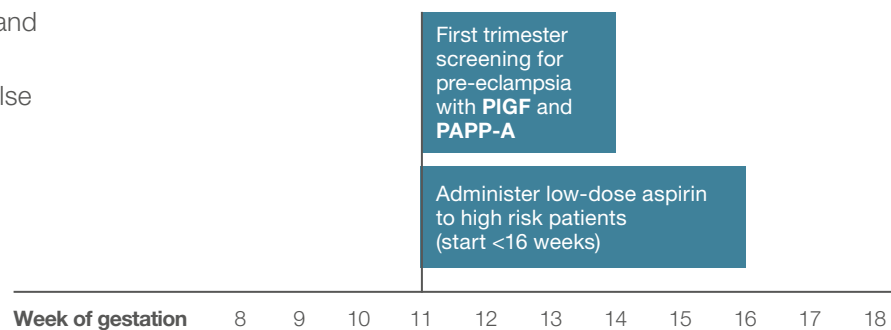
Combined screening for pre-eclampsia in weeks 11–13+6 can reliably identify women at risk for developing pre-eclampsia.

Combined first trimester screening includes

- serum PIGF and PAPP-A measurement,
- determination of mean arterial pressure (MAP), and
- Uterine Artery Pulsatility Index (UAPI)

resulting in a detection rate of >90% for a fixed false positive rate of 5%.<sup>17</sup>

An early identification of high-risk women allows for preventive measures and intensified monitoring. Administering low-dose aspirin (<150 mg/day) to high-risk women before 16 weeks of gestation can significantly reduce the incidence of pre-eclampsia by 50%–90%.<sup>18,19</sup>



### Facts on pre-eclampsia

- Multisystem, life-threatening pregnancy-related disorder
- A main reason for maternal and fetal morbidity and mortality<sup>20,21</sup>
- **Incidence:** 2-8% of pregnancies
- **Definition:** New onset hypertension and proteinuria >20 weeks of gestation in previously normotensive women<sup>22</sup>
- **HELLP** (Hemolysis, Elevated Liver enzymes, Low Platelets): Severe pre-eclampsia variant occurring in ≈20% of symptomatic women; defined by additional affection of liver and coagulation system<sup>23</sup>
- **Eclampsia:** Final stage of disease, associated with severe tonic-clonic seizures and coma as well as brain injury, cerebral edema and stroke<sup>23</sup>



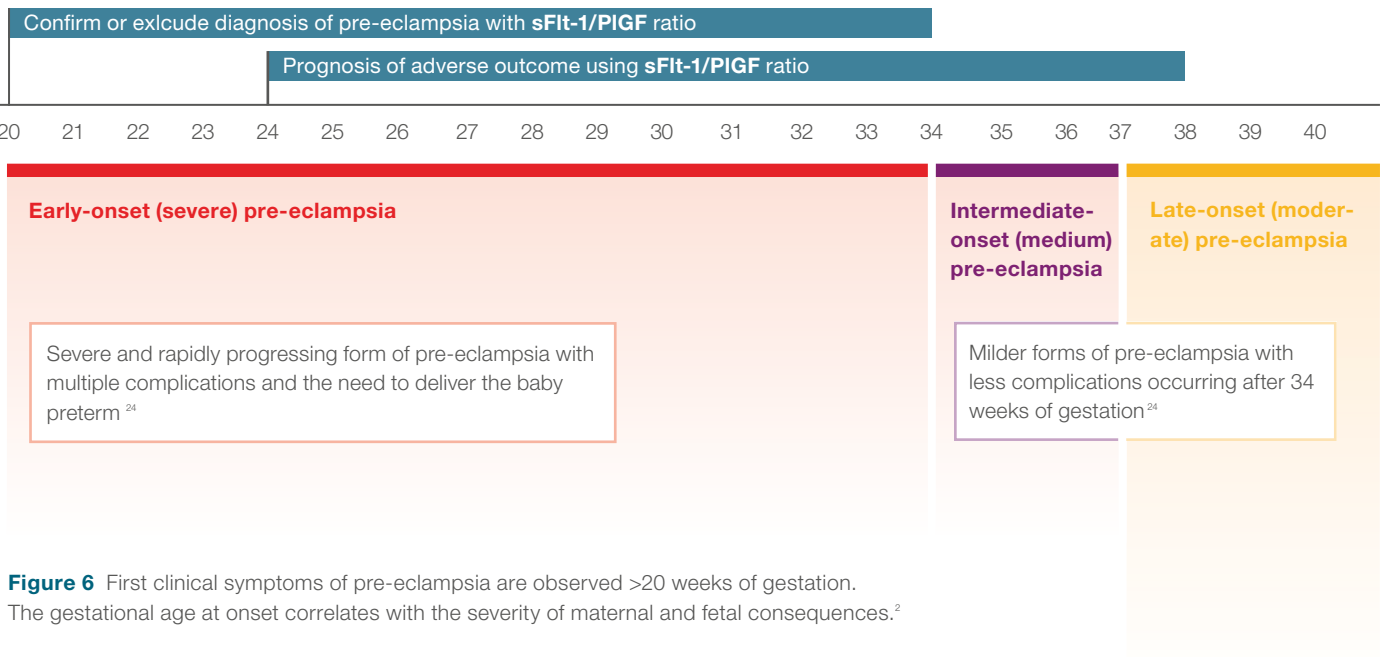
## sFlt-1/PlGF ratio: Improved diagnosis and prognosis of adverse outcome

First symptoms of pre-eclampsia (hypertension, proteinuria) are observed after 20 weeks of gestation.<sup>23</sup>

Diagnosis of pre-eclampsia is difficult, as pre-eclampsia can be confused with other diseases such as pregnancy-induced hypertension.

By adding sFlt-1/PlGF ratio to the current diagnostic standard, the **diagnosis of pre-eclampsia** in a symptomatic woman can be confirmed or excluded.<sup>2,10,11</sup>

In women with diagnosed pre-eclampsia, the sFlt-1/PlGF ratio is a potent **predictor of subsequent maternal and fetal adverse outcome** and can be useful for further patient management.<sup>5,6</sup>



**Figure 6** First clinical symptoms of pre-eclampsia are observed >20 weeks of gestation. The gestational age at onset correlates with the severity of maternal and fetal consequences.<sup>2</sup>



# Complete pre-eclampsia portfolio

From safe screening to improved diagnosis  
with B·R·A·H·M·S sFlt-1 and PIGF plus

## Thermo Scientific B·R·A·H·M·S sFlt-1 KRYPTOR

Automated immunofluorescent assay for the quantitative determination of the concentration of sFlt-1 (soluble FMS-like Tyrosine Kinase 1, also known as VEGF receptor-1) in human serum.

- 75 determinations per kit
- 9 min incubation time
- Monoparametric control kit, 3 levels
- Wide measuring range: 22-90000 pg/mL
- Excellent precision

With the lower and upper detection limits of 22 and 90000 pg/mL B·R·A·H·M·S sFlt-1 KRYPTOR provides the measuring range needed for a **reliable detection of clinical sFlt-1 values throughout pregnancy.**



sFlt-1  
PIGF



**Thermo Scientific**  
**B·R·A·H·M·S PIGF plus KRYPTOR**

Automated immunofluorescent assay for the quantitative determination of the concentration of PIGF (Placental Growth Factor) in human serum. The assay is specific for the measurement of human free PIGF-1.

- **75 determinations per kit**
- **29 min incubation time**
- **Monoparametric control kit, 3 levels**
- **Wide measuring range: 3.6-7000 pg/mL**
- **Excellent precision**

With a detection limit of 3.6 pg/mL and an upper limit of 7000 pg/mL B·R·A·H·M·S PIGF plus KRYPTOR provides the high sensitivity needed for **measuring PIGF levels in first trimester** as well as a wide measuring range to **reliably measure clinical values in second and third trimester**.

Exceptionally precise, fast and easy

**Thermo Scientific B·R·A·H·M·S KRYPTOR compact PLUS**

18 Years Reliable Results

18 Years Confident Decisions

- **All KRYPTOR platforms FMF approved**
- **In routine use by FMF since 1999**
- **Excellent precision and proven median stability**

## Your **BENEFITS** by adding the sFit-1/PIGF ratio into clinical routine

- ▶ **Improved clinical accuracy** in diagnosing pre-eclampsia in symptomatic patients
- ▶ **Potent prognostic tool** for subsequent adverse pregnancy outcome

## Your **ACCESS** to our interactive e-detail

Get more information on pre-eclampsia management throughout pregnancy:



<http://prenatal.world-of-biomarkers.com>

Pin code: **ratio01**



### Thermo Scientific B·R·A·H·M·S Biomarkers Prenatal Screening Portfolio on KRYPTOR Systems

B·R·A·H·M·S <b>AFP</b> KRYPTOR	Art. no.: 816.075
B·R·A·H·M·S <b>Free <math>\beta</math>hCG</b> KRYPTOR	Art. no.: 809.075
B·R·A·H·M·S <b>hCG+<math>\beta</math></b> KRYPTOR	Art. no.: 841.050
B·R·A·H·M·S <b>Inhibin A</b> KRYPTOR	(under development)
B·R·A·H·M·S <b>PAPP-A</b> KRYPTOR	Art. no.: 866.075
B·R·A·H·M·S <b>PIGF plus</b> KRYPTOR*	Art. no.: 859.075
B·R·A·H·M·S <b>sFit-1</b> KRYPTOR*	Art. no.: 845.075
B·R·A·H·M·S <b>uE3</b> KRYPTOR**	Art. no.: 803.075
B·R·A·H·M·S <b>Fast Screen pre I plus Software</b>	Art. no.: 105750

\* Available on KRYPTOR compact PLUS

\*\* Available on KRYPTOR and KRYPTOR compact PLUS

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