



STANDING COMMITTEE  
FOR QUALITY AND COMPETENCE (QCC)



With the financial support from the Prevention of and Fight against Crime Programme of the European Union  
European Commission – Directorate-General Home Affairs

# Guidelines for the single laboratory Validation of Instrumental and Human Based Methods in Forensic Science **Examples**

DOCUMENT TYPE:	REF. CODE:	ISSUE NO:	ISSUE DATE:
GUIDANCE - ANNEX	QCC-VAL-002	001	10/11/2014

## Project Team

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## 5 Instrument-based examples

### 5.1 Primarily quantitative

#### 5.1.1 Quantification of DNA using Real-Time PCR

## SINGLE LABORATORY VALIDATION OF MEASUREMENT PROCEDURES (Quantification System)

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**Sample validation-Guideline purposes only**

**1. Specify the measurement procedure, analyte, measurand and units**

<i>The measurement procedure</i>	Quantitative analysis of DNA samples using (1) Quantifiler™ Human DNA and (2) Quantifiler™ Y Human DNA kits
<i>Analyte</i>	Quantifiler™ Human DNA- Total human genomic DNA (male and female)
<i>The measurand</i>	Concentration of human genomic DNA and concentration of male genomic DNA in casework samples
<i>Unit</i>	ng/ul of human genomic DNA in extract ng/ul of male genomic DNA in extract

**2. Specify the Scope**

<i>Matrix</i>	Case samples
<i>Measuring range</i>	50ng/ul-0.023ng/ul (23pg)

**3. Requirement on the measurement procedure**

<i>Intended use of the results</i>	<p>The Quantifiler™ Human DNA assay is intended for general use as a preliminary DNA quantification step to performing routine STR analysis</p> <p>The Human Male kit assay is designed to be of use particularly in samples with mixed male-female DNAs, such as sexual assault evidence, where it may be useful to separately detect and quantify male DNA from a background of female DNA.</p>
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**Internal validation:** To ensure kit performs as expected in the laboratory.

<i>Mark the laboratory/customer requirements and give their values</i>	<i>Parameters to be validated</i>		<i>Value requested</i>
	<i>x</i>	Known and case samples	<p>% Difference from Quantifier Human kit and QuantiBlot (system Quantifiler replacing) should not be significant.</p> <p>Using quantification values of 40pg/ul to 2ng/ul for the Quantifier kits, complete profiles for all samples, with no off-scale or drop-out allele peaks should be obtained This would not be achievable in the case of inaccurate DNA quantification.</p>

<b>x</b>	Precision	<p>The mean <math>C_{T\text{FAM}}</math> results for the DNA standards should show the normal inverse relationship with sample DNA concentration, going from the range of approximately 23 to 25 for the highest standard (50 ng/<math>\mu\text{L}</math>) to 33 to 36 for the lowest standard (23 pg/<math>\mu\text{L}</math>). A linear relationship between the <math>C_{T\text{FAM}}</math> (template DNA) and concentration should be obtained.</p> <p>Range of the <math>C_{T\text{Vic}}</math> (IPC-internal PCR control) in the range of approximately 27 to 29 for all standards and samples.</p>
<b>x</b>	Reproducibility	<p>Reproducible <math>C_{T\text{Vic}}</math> values for the IPC and <math>C_{T\text{FAM}}</math> values should be obtained in replicates, between operators and on using two 7500 real time PCR instruments</p> <p>Reproducible DNA profile quality as assessed by determining average peak heights should be obtained.</p>
<b>x</b>	Sensitivity /Stochastic	Quantification result obtained at 0.023ng/ul (23pg)
<b>x</b>	Mixture Studies	<p>Quantifier Human kit should show the expected increase in total human genomic DNA as a result of adding increasing concentrations of K562 (female DNA)</p> <p>Quantifier Y should remain unaffected by the increase in female DNA and detect only the constant level of male DNA.</p>
<b>x</b>	False Positive rate	Quantifier kit assays prepared in the absence of DNA give a $C_{T\text{FAM}} > 50$

#### 4. Origin of the Measurement Procedure

		VALIDATION
<i>New In-House Method</i>	<b>X</b>	<i>Full internal validation</i>
<i>Modified Validated Method</i>	<input type="checkbox"/>	<i>Partial</i>

<i>Official Standard Method</i>	<input type="checkbox"/>	<i>“Simple”</i>
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## 5. Performance Parameters

Parameter	Description	
Known and case samples	<p>Quantifiler™ Human DNA and Quantifiler™ Y Human DNA kits must be evaluated and tested using known and case samples (actual and mock) and compared to previous quantification method(s).</p>	
	<p style="text-align: center;"><b>Demand</b></p>	
	<p>% difference from Quantifier Human kit and QuantiBlot should not be significant.</p> <p>For SGM+ STR assays, set up with DNA input amounts determined by the results of Quantifiler kit assays to provide complete profiles for all samples, with no off-scale or drop-out allele peaks that would have resulted from inaccurate DNA quantification.</p>	
	<p style="text-align: center;"><b>Experiment</b></p>	
	<p>Fifty reference samples (taken from laboratory staff- 25 male and 25 female) and thirty casework samples were quantitated with Quantifier kits and QuantiBlot, amplified with SGM+ and analysed on a 3100 Genetic Analyzer. It was deemed that the range of quantification data would be covered with these samples.</p> <p>SGM+ STR reactions were adjusted to give a more consistent amount of 1ng/ul if the DNA concentration was high enough. The samples with a low concentration were added to the SGM+ reactions at the maximum allowed volume of 20ul. SGM+ kit assay results were then obtained by electrophoresis on the ABI PRISM 3100 genetic analyser, and assessed according to the criterion that STR peaks should be within the range to generate successful genotype results (i.e. no drop out or off scale peaks)</p>	
	<p style="text-align: center;"><b>Evaluation</b></p>	
	<p>% Differences were calculated as Quantifiler Human kit results/QuantiBlot results DNA profiles quantitated with Quantifier kits assessed over the range of concentrations 10pg/ul and 2.0ng/ul.</p>	
<p style="text-align: center;"><b>Results:</b></p>	<p>% Differences between the Quantifier Human kit and QuantiBlot systems exceeded 50%.</p> <p>The difference can for the most part be explained because the comparison in the QuantiBlot system is done by a visual comparison against the DNA standards as against extrapolation from a standard curve. A more accurate comparison of % differences would be achievable if comparing two equivalent DNA standard curve systems. As the Quantifier Human kit gave overall higher quantification data it was deemed to be more fit for purpose than the existing technique of QuantiBlot.</p> <p>Concordant and successful genotypes with no drop out or off scale peaks obtained for Quantifier systems over the range of concentrations 40pg/ul and 2.0ng/ul. Profiles were also obtained for</p>	
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	20pg/ul (.2ng/ul in 20ul) but imbalance was observed in the peaks
<b>Conclusions:</b>	Fit for purpose

<b>Parameter</b>	<b>Description</b>
<b>Precision</b>	The tightness of the quantification results
	<b>Demand</b>
	The mean $C_{T\text{FAM}}$ results for the DNA standards going from the range of approximately 23 to 25 for the highest standard (50 ng/ $\mu\text{L}$ ) to approximately 33 to 36 for the lowest standard (23 pg/ $\mu\text{L}$ ). Range of the $C_{T\text{Vic}}$ (IPC) in the range of approximately 27 to 29 for all standards and samples.
	<b>Experiment</b>
	A set of 8 standard dilutions of Quantifiler DNA standards were made ranging in concentrations of 50ng to 0.023ng. These were run in 3 separate plates on 2 separate days.  Twenty single source samples were quantified on three different days. Each of the twenty samples was also quantified in triplicate on a single run. Male: female mixtures were also prepared and quantified in triplicate over several days
	<b>Evaluation</b>
	The $C_{T\text{FAM}}$ values were compiled, averaged and SD determined. Average $C_{T\text{FAM}}$ values for 8 DNA standards were plotted against concentration to ensure linearity  Also, the $C_{T\text{Vic}}$ values were compiled, averaged and SD determined

<b>Results:</b>	<p>The mean <math>C_{T\text{FAM}}</math> results for the DNA standards going from the range of 23 to 25 for the highest standard (50 ng/<math>\mu\text{L}</math>) to 33 to 36 for the lowest standard (23 pg/<math>\mu\text{L}</math>).</p> <p>Linearity achieved.</p> <p>Range of the <math>C_{T\text{VIC}}</math> (IPC) 27 to 29 for all standards and samples</p>
<b>Conclusions:</b>	Fit for purpose

Parameter	Description
<b>Reproducibility</b>	The ability to obtain the same results under the same conditions
	<b>Demand</b>
	Reproducibility of $C_{T\ vic}$ values for the IPC and $C_{T\ FAM}$ for the template DNA Reproducibility in average peak heights obtained
	<b>Experiment</b>
	Twenty single source samples were quantified on three different days. Each of the twenty samples was also quantified in triplicate on a single run. Male: female mixtures were also prepared and quantified in triplicate over several days ( <b>same samples as precision samples</b> )
	A sample of K562 (cell line DNA-female) was diluted from 2ng/ul to 0.06ng/ul and quantitated in replicates of 4 by two separate analysts on two separate days for at least three runs. Two 7500 RT-PCR instruments used
	<b>Evaluation</b>
The $C_{T\ vic}$ values for the IPC were compiled The $C_{T\ FAM}$ values for the template DNA were compiled  Samples were selected from the reproducibility study, amplified using SGM+, run on 3100 genetic analyzer and the average peak heights determined	
<b>Results:</b>	Reproducible $C_{T\ vic}$ values for the IPC and $C_{T\ FAM}$ values for the template DNA obtained in replicates, between operators and on using two 7500 instruments over a range of concentrations of DNA  Reproducibility in average peak heights obtained
<b>Conclusions:</b>	Fit for purpose



Parameter	Description
<b>Sensitivity</b>	Studies to determine sensitivity
	<b>Demand</b>
	Quantification result obtained at 0.023ng/ul (23pg)
	<b>Experiment</b>
	Series of dilutions 1/10 to 1/1000 were made using a male DNA control sample (5ng/ul) in Tris EDTA buffer. The dilutions were run in triplicate on 3 different plates.
	<b>Evaluation</b>
	DNA quantification data obtained for the dilutions assessed
<b>Results:</b>	Quantifier kits could detect DNA in 1/1000 dilution (5pg/ul). While DNA can be detected at these levels poor profile quality/no profile was obtained at these low levels of DNA.
<b>Conclusions:</b>	Fit for purpose

Parameter	Description
<b>Mixtures studies</b>	Quantifier reaction should show the large range of concentrations in the different mixture samples and distinguish different contributors (male/female).
	<b>Demand</b>
	Lowest amount of male DNA that can be still amplified and detected in the presence of female DNA (total) (0.05ng/ul male detected in presence of 50ng/ul of female DNA)
	<b>Experiment</b>
	A series of mixtures were made 1:1, 1:5, 1:25, 1:50, 1:100, 1:250, 1:500 and 1:1000 using a mixture of male and female DNA samples in a final volume of 100ul. Male DNA sample- Quantifier DNA standard in kit (200ng/ul) Female DNA sample used –K562 DNA standard, 623ng/ul. The male DNA was kept constant in all dilutions at a concentration of 50pg/ul (0.05ng/ul). The female DNA ranged from 50pg/ul in the 1:1 dilution to 50ng/ul in the 1:1000 dilution. Each sample was analysed in three replicates reactions each with the Quantifier Human and Quantifier Y assays.
<b>Evaluation</b>	
Results from the three replicates reactions each with the Quantifier Human and Quantifier Y assays were evaluated to determine if (1) the Quantifier Human showed the expected increase in total human genomic DNA as a result of the adding K562 and (2) the Quantifier Y remained un affected by the increase in female DNA detecting only the constant level of male DNA.	
<b>Results:</b>	Quantifier Human showed the expected increase in total human genomic DNA as a result of the adding K562 Quantifier Y remained un affected by the increase in female DNA detecting only the constant level of male DNA.
<b>Conclusions:</b>	Fit for purpose

Parameter	Description
<b>False positive rate</b>	Assessment of false positive rate
	<b>Demand</b>
	Quantifier kit assays prepared in the absence of DNA give a $CT_{FAM} > 50$
	<b>Experiment</b>
	A set of 50 reactions of each Quantifiler kit assay (Human and Y) were prepared and dispensed as usual. Each reaction received 2 $\mu$ L of TE Buffer as its sample volume. The assays were run as per the standard protocol except for one change: thermal cycling was extended to 50 cycles. The samples were analyzed with the SDS software after completion of the runs, and those with a CT that was $< 50$ would have been considered to be false-positive results.
	10 reagent blanks from previous proficiency trials were quantified
<b>Evaluation</b>	
The samples were analyzed after completion of the runs, and those with a CT that was $< 50$ would have been considered to be false-positive results.	
<b>Results:</b>	All samples gave CT that was $> 50$
<b>Conclusions:</b>	Fit for purpose

## 6. Conclusions

<b>Summary</b>	All parameters tested deemed fit for purpose and no additional work required.	
<b>Plan of internal quality control:</b>	<p>-Quality assurance parameters and reporting guidelines derived from internal validation results</p> <p><b>-For calibration curve</b></p> <p>An <math>R^2</math> value <math>\geq 0.99</math> acceptable (closeness of fit between the standard curve regression line and the individual <math>C_{T\text{FAM}}</math> data points of Quantification standard reactions)</p> <p>-Slope indicates the PCR amplification efficiency. For Quantifiler™ Human DNA slope values required between <math>-2.9</math> and <math>-3.3</math></p> <p>For Quantifiler™ Y DNA slope values required between <math>-3.0</math> and <math>-3.6</math></p> <p>-<math>C_{T\text{FAM}}</math> results for the DNA standards used to monitor the quality of batch lots of quantification kits</p>	
<b>Statement on fitness for intended purpose</b>	<input checked="" type="checkbox"/> <b>The method fulfils the demand</b>	<input type="checkbox"/> <b>The method does not fulfil the demand</b> <b>The following should be done....</b>

### Reference Documents

SWGDM Validation guidelines for DNA Analysis Methods  
Developmental Validation of the Quantifiler™ Real-Time PCR Kits for the Quantification of Human Nuclear DNA Samples. *J Forensic Sci*, July 2005, Vol. 50, No. 4

## 5.1.2 Quantification of alcohol in blood

### SINGLE-LABORATORY VALIDATION OF MEASUREMENT PROCEDURES

#### 1. Specify the measurement procedure, analyte, measurand and units









<i>The measurement procedure</i>	Quantitative analysis of ethanol in blood and urine using headspace GC-FID
<i>Analyte</i>	Ethanol
<i>The measurand</i>	Concentration of ethanol in liquid
<i>Unit</i>	g/L

#### 2. Specify the Scope


<i>Sample type – matrix</i>	Human blood samples
<i>Measuring range</i>	0.1 – 3 g/L
<i>Intended use of the results</i>	Court Reports and Intelligence reports to Police force

#### 3. Requirement on the measurement procedure

Remark: the criteria set for precision and bias are the legal requirements in Belgium.

<i>Parameters to be validated</i>		<i>Value requested</i>
<i>Precision</i>	<i>Repeatability</i>	 RSD <12% (0.2 g/L) RSC <8% (>0.4 g/L)
	<i>Within-lab reproducibility</i>	 RSD <12% (0.2 g/L) RSD <8% (>0.4 g/L)
<i>Trueness</i>	<i>Test for bias</i>	 Bias <15 % (0.1 0.4 g/L) Bias < 5% (0.41 – 0.69 g/L) Bias <3% (>0.7 g/L)
	<i>Selectivity</i>	 Test for possible interference Distinguishability of compounds and the internal control
<i>Measurement range</i>	<i>LOD</i>	 0.025 g/L
	<i>LOQ</i>	 0.1 g/L
	<i>Linearity</i>	 Residual < 3 % (> 0,4l) Residual < 0,01 (0,4 g/l)
<i>Ruggedness (Robustness)</i>		Investigation towards influence of the matrix
<i>Stability</i>	<i>Processed samples</i> <i>Freeze-thaw</i> <i>Longterm storage</i>	 stable

#### 4. Origin of the Measurement Procedure

		VALIDATION
<i>New In-House Method</i>		<i>Full</i>
<i>Modified Validated Method</i>		<i>Partial</i>
<i>Official Standard Method</i>		“Simple”

Parameter	Description
<b>Precision</b>	Repeatability, Within-Lab Reproducibility,
	<b>Demand</b>
	Repeatability/Within-Lab Reproducibility for low amounts (0.2 g/L) lower than 12% Repeatability/Within-Lab Reproducibility for higher amounts (>0.4 g/L) lower than 8%
	<b>Experiment</b>
	9 control samples (6 in serum and 3 in full blood) were processed in duplo, and this at 8 different days.
	Results of the measurements with serum control samples.

Serum controls Conc. (g/L)	Run1	Run2	Run3	Run4	Run5	Run6	Run7	Run8
0,2	0,212	0,199	0,206	0,206	0,206	0,213	0,210	0,206
	0,209	0,198	0,203	0,204	0,199	0,207	0,214	0,209
0,5	0,499	0,516	0,499	0,513	0,506	0,548	0,516	0,520
	0,497	0,514	0,508	0,510	0,495	0,505	0,506	0,526
0,8	0,796	0,782	0,793	0,793	0,794	0,800	0,795	0,802
	0,788	0,783	0,790	0,798	0,791	0,794	0,806	0,797
1,1	1,128	1,124	1,120	1,146	1,119	1,108	1,129	1,135
	1,122	1,135	1,162	1,125	1,111	1,128	1,133	1,146
1,5	1,475	1,489	1,520	1,503	1,513	1,502	1,468	1,550
	1,490	1,493	1,506	1,524	1,494	1,472	1,481	1,482
3	3,048	3,039	3,132	3,050	3,119	2,989	2,988	3,048
	2,955	3,020	3,145	3,109	3,132	3,168	3,081	3,072

Calculations demonstrated for serum control 0.2g/L

Anova: Single Factor

SUMMARY

Groups	Count	Sum	Average	Variance
Column 1	2	0,421049	0,2105245	2,98901E-06
Column 2	2	0,397517	0,1987585	1,00962E-06
Column 3	2	0,408793	0,2043965	5,89961E-06
Column 4	2	0,409996	0,204998	4,2108E-06
Column 5	2	0,405196	0,202598	2,47526E-05
Column 6	2	0,420174	0,210087	1,87762E-05
Column 7	2	0,424031	0,2120155	8,88311E-06
Column 8	2	0,41456	0,20728	4,15873E-06

ANOVA

Source of Variation	SS	df	MS	F	P-value	F crit
Between Groups	0,000283409	7	4,0487E-05	4,582581513	0,024177467	3,500463855
Within Groups	7,06797E-05	8	8,83497E-06			
Total	0,000354088	15				

Repeatability:

$$s_r^2 = MS_{\text{within}} = \text{Average of all measurements} = 8,83497E-06$$

$$So: RSD_r [\%] = 0,206$$

$$1,44$$

Bias [%]:

3,68

Time-different intermediate precision:

$$s_t^2 = (MS_{\text{between}} - MS_{\text{within}})/n = 1,5826E-05$$

$$s_t^2 + s_r^2 = 2,4661E-05$$

$$So: RSD_{(t)} [\%] = 2,41$$

Results:

Control sample	Repeatability (%)	Within-lab reproducibility (%)
0.2 g/L serum	1.44	2.41
0.5 g/L serum	2.3	2.59
0.8 g/L serum	0.52	0.82
1.1 g/L serum	1.21	1.21

	1.5 g/L serum	1.39	1.45
	3.0 g/L serum	1.89	2.04
	0.5 g/L full blood	1.09	1.54
	0.8 g/L full blood	1.58	1.73
	1.1 g/L full blood	1.66	2.08
<b>Conclusions:</b>	Fit for purpose		

Parameter	Description																				
<b>Bias</b>	Laboratory bias																				
	Selectivity																				
	<b>Demand</b>																				
	Depending on the amount of alcohol in the sample: <15% for low concentrations (0.1-0.4 g/L) <5 % for moderate concentrations (0.41 – 0.69 g/L) <3% for higher concentrations (<0.7 g/L)																				
	<b>Experiment</b>																				
	Bias calculation included in within-lab reproducibility analysis.  Selectivity: Determination if the following compounds can be discriminated from ethanol: methanol, acetone, isopropanol, n-propanol, t-butanol.																				
<b>Results:</b>	<table border="1"> <thead> <tr> <th>Control sample</th> <th>Bias (%)</th> </tr> </thead> <tbody> <tr> <td>0.2 g/L serum</td> <td>3.68</td> </tr> <tr> <td>0.5 g/L serum</td> <td>-0.35</td> </tr> <tr> <td>0.8 g/L serum</td> <td>-1.98</td> </tr> <tr> <td>1.1 g/L serum</td> <td>-1.61</td> </tr> <tr> <td>1.5 g/L serum</td> <td>0.04</td> </tr> <tr> <td>3.0 g/L serum</td> <td>0.80</td> </tr> <tr> <td>0.5 g/L full blood</td> <td>0.97</td> </tr> <tr> <td>0.8 g/L full blood</td> <td>0.64</td> </tr> <tr> <td>1.1 g/L full blood</td> <td>0.77</td> </tr> </tbody> </table>	Control sample	Bias (%)	0.2 g/L serum	3.68	0.5 g/L serum	-0.35	0.8 g/L serum	-1.98	1.1 g/L serum	-1.61	1.5 g/L serum	0.04	3.0 g/L serum	0.80	0.5 g/L full blood	0.97	0.8 g/L full blood	0.64	1.1 g/L full blood	0.77
	Control sample	Bias (%)																			
	0.2 g/L serum	3.68																			
	0.5 g/L serum	-0.35																			
	0.8 g/L serum	-1.98																			
	1.1 g/L serum	-1.61																			
	1.5 g/L serum	0.04																			
	3.0 g/L serum	0.80																			
	0.5 g/L full blood	0.97																			
	0.8 g/L full blood	0.64																			
	1.1 g/L full blood	0.77																			
Selectivity: All compounds can be separated except acetone and t-butanol.																					
<b>Conclusions:</b>	Fit for purpose																				



Parameter	Description																										
<b>Measuring range</b>	<ul style="list-style-type: none"> <li>• Linearity</li> <li>• Limit of Quantification (LOQ)</li> <li>• Limit of Detection (LOD)</li> </ul>																										
	<b>Demand</b>																										
	<ul style="list-style-type: none"> <li>• Residual &lt; 3% (&lt;0.4 g/L) Residual &lt; 0.01 (0.4g/L)</li> <li>• Which is the lowest concentration still to be quantified</li> <li>• Which is the lowest concentration still to be detected in a reproducible manner</li> </ul>																										
	<b>Experiment</b>																										
	<ul style="list-style-type: none"> <li>• Linearity Ethanol calibration curves were set up using standards for 6 times and the Residual % was calculated</li> <li>• Limit of quantification: <ul style="list-style-type: none"> <li>a. By testing A 0.1 g/L control sample in either serum or full blood was run during 8 days in duplo. The bias, repeatability and within-lab reproducibility was calculated and verified if this is still within criteria.</li> <li>b. By calculation. Following the above experiment the LOQ can be calculated as 10sRw</li> </ul> </li> <li>• Limit of detection. A dilution series was set up of a control sample in serum and in full blood. The signal to noise ratio were determined for each sample in a duplo analysis. S/N values above 5 were determined as positive.</li> </ul>																										
	<b>Evaluation</b>																										
<p>Linearity</p> <p>Limit of Quantification:</p> <p>a. Determination by testing (control sample as 0.1 g/L)</p> <table border="1"> <thead> <tr> <th>Matrix</th> <th>Bias (%)</th> <th>Repeatability (%)</th> <th>Within-lab reproducibility (%)</th> </tr> </thead> <tbody> <tr> <td>Serum</td> <td>9.08</td> <td>3.06</td> <td>5.1</td> </tr> <tr> <td>Full blood</td> <td>9.31</td> <td>1.63</td> <td>3.12</td> </tr> </tbody> </table> <p>b. determination by calculation (LOQ = 10s : sRw) LOQ in serum: 0.05 g/L LOQ in full blood: 0.03 g/L</p> <p>Limit of detection</p> <table border="1"> <thead> <tr> <th>Dilution to concentration</th> <th>S/N ratio in serum</th> <th>S/N ratio in blood</th> </tr> </thead> <tbody> <tr> <td>0.05</td> <td>13.5</td> <td>10.2</td> </tr> <tr> <td>0.025</td> <td>5.65</td> <td>5.4</td> </tr> <tr> <td>0.01</td> <td>2.55</td> <td>2.3</td> </tr> <tr> <td>0.005</td> <td>1.25</td> <td>1.25</td> </tr> </tbody> </table>	Matrix	Bias (%)	Repeatability (%)	Within-lab reproducibility (%)	Serum	9.08	3.06	5.1	Full blood	9.31	1.63	3.12	Dilution to concentration	S/N ratio in serum	S/N ratio in blood	0.05	13.5	10.2	0.025	5.65	5.4	0.01	2.55	2.3	0.005	1.25	1.25
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<b>Results:</b>	<p>Linearity: Residual% &lt;3% (&gt;0.4g/L) Residual &lt;0.01 (0.4g/L)</p> <p>The limit of quantification: the 0.1 g/L sample gave a reproducible and correct result. The calculation following these measurements predicted the real LOQ at 0.05 g/L in serum and 0.03 g/L in full blood.</p> <p>The limit of detection: the dilution sample to concentration of 0.025 %gave an average S/N value of &gt;5</p>
<b>Conclusions:</b>	Fit for purpose

Parameter	Description
<b>stability</b>	Stability
	<b>Demand</b>
	Measure the stability of results <sup>1</sup> I. Post-processed in the autosampler (room temperature 24h) II. Freeze-thaw (3 cycli) III. Long-term storage 4°C / -20°C
	<b>Experiment</b>
	I. QC samples (0.5 g/L and 1.1 g/L in full blood) were prepared, and analyses freshly (set a) or after storage 24 hours in the autosamples (set b) II. QC samples (0.5 g/L and 1.1g/L in full blood) were prepared and freeze-thawed three times. The stability samples (set b) were processed and compared to QC samples prepared freshly at the day of the process as control samples (set a) III. QC samples (0.5 g/L and 1.1 g/L in full blood) were prepared and stored for a longer period (2 months) at respectively 4°C or -20°C. The stability samples (set b) were processed and compared to QC samples prepared freshly at the day of the process as control samples (set a)  Comparison is done by comparing the average peak areas, the ratio (b/a), and the determination of the 90% confidence interval and 80-120% acceptance interval.
	<b>Evaluation</b>

<sup>1</sup> Based on Wille et al. (2011)

**A. Stability (24h) of processed samples in autosampler**

QC sample	0.5 g/L	1.1 g/L
Average area ratio (ethanol/n-propanol) of the control samples (a)	0.322	0.716
Average area ratio (ethanol/n-propanol) of the stability samples (b)	0.328	0.704
Ratio b/a*100%	101.8%	98.5
90% Confidence interval of the area ratio of the stability samples	0.324-0.332	0.697-0.711
80-120% Acceptance interval of the area ratio of the control samples	0.258-0.387	0.572-0.859

**B. Stability (freeze-thaw) of processed samples**

QC sample	0.5 g/L	1.1 g/L
Average area ratio (ethanol/n-propanol) of the control samples (a)	0.319	0.722
Average area ratio (ethanol/n-propanol) of the stability samples (b)	0.339	0.758
Ratio b/a*100%	106.2%	105.1%
90% Confidence interval of the area ratio of the stability samples	0.333-0.345	0.752-0.764
80-120% Acceptance interval of the area ratio of the control samples	0.255-0.383	0.577-0.866

**Ca. long-term storage 4°C**


QC sample	0.5 g/L	1.1 g/L
Average area ratio (ethanol/n-propanol) of the control samples (a)	0.322	0.716
Average area ratio (ethanol/n-propanol) of the stability samples (b)	0.312	0.720
Ratio b/a*100%	96.8%	100.6%
90% Confidence interval of the area ratio of the stability samples	0.310-0.314	0.698-0.742
80-120% Acceptance interval of the area ratio of the control samples	0.258-0.387	0.572-0.859

**Cb. Long-term storage -20°C**

QC sample	0.5 g/L	1.1 g/L
Average area ratio (ethanol/n-propanol) of the control samples (a)	0.322	0.716
Average area ratio (ethanol/n-propanol) of the stability samples (b)	0.310	0.714
Ratio b/a*100%	96.3%	99.8%
90% Confidence interval of the area ratio of the stability samples	0.295-0.325	0.700-0.728
80-120% Acceptance interval of the area ratio of the control samples	0.258-0.387	0.572-0.859

<b>Results:</b>	No significant change in values is obtained by any kind of stability study
<b>Conclusions:</b>	Fit for purpose

## 6. Conclusions

<b>Summary</b>	Laboratory/customer demands met	
<b>Plan of internal quality control:</b>	Control charts put in place to ensure the system is operating within its specifications.	
<b>Statement on fitness for intended purpose</b>	 <b>The method fulfills the demand</b>	<input type="checkbox"/> <b>The method does not fulfill the demand</b> <b>The following should be done....</b>

### Reference Documents

1. Wille, S., F. Peters, V. Di Fazio & N. Samyn. Practical aspects concerning validation and quality control for forensic and bioanalytical quantitative methods. 2011. Accred Qual Assur 16: 279-92.

### 5.1.3 Quantification of cocaine in powder

## SINGLE-LABORATORY VALIDATION OF MEASUREMENT PROCEDURES

Sample validation-Guideline purposes only








### 1. Specify the measurement procedure, analyte, measurand and units

<i>The measurement procedure</i>	Quantitative analysis of cocaine using GC-FID
<i>Analyte</i>	Cocaine
<i>The measurand</i>	Concentration of cocaine in powder
<i>Unit</i>	% (w/w)


### 2. Specify the Scope

<i>Sample type – matrix</i>	Street drugs samples delivered to the laboratory
<i>Measuring range</i>	1 - 100 weight%
<i>Intended use of the results</i>	Court Reports and Intelligence reports to Police Force  Note: Before any sample is submitted for quantitative analysis it will already have been analysed qualitatively and identified by GCMS. This is a quantitative method with a low demand on uncertainty

### 3. Requirement on the measurement procedure

<i>Parameters to be validated</i>		<i>Value requested</i>
<i>Precision</i>	<i>Repeatability</i>	 RSD <sub>r</sub> 5 %
	<i>Within-lab reproducibility</i>	 RSD <sub>Rw</sub> 7%
<i>Trueness</i>	<i>Test for bias</i>	 Bias < 10 % relative
	<i>Selectivity</i>	 Test for possible interference
<i>Measurement range</i>	<i>LOD or LOQ</i>	 LOQ 1 %
	<i>Linearity</i>	 Residual max deviation 2 % relative
<i>Ruggedness (Robustness)</i>		Routine GC method – not necessary
<i>Measurement uncertainty</i>		 Expanded uncertainty < 20 % relative

### 4. Origin of the Measurement Procedure

		VALIDATION
<i>New In-House Method</i>		<i>Full</i>
<i>Modified Validated Method</i>		<i>Partial</i>
<i>Official Standard Method</i>		<i>“Simple”</i>



Parameter	Description																																																																																																																																																			
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	RSD <sub>Rw</sub> - 3 Analysts, 3 analysis (in duplicate) per day on different days of check standard cocaine sample																																																																																																																																																			
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Column C- Wt of Cocaine (3-4mg)																																																																																																																																																				
Column D-purity of Cocaine																																																																																																																																																				

***S<sub>Rw</sub>* Results obtained**

	<i>B</i>	<i>C</i>	<i>D</i>		Analyst 1	Analyst 2	Analyst 3		
H1502006	13.48	3.601	23.86						
H1502008	13.48	3.579	23.71	23.78	Sample 1	23.78	24.54	24.42	
H1502020	11.81	3.235	24.46		Sample 2	24.43	24.85	25.10	
H1502021	11.81	3.227	24.40	24.43	Sample 3	24.23	24.18	24.84	
H1502032	13.15	3.556	24.15						
H1502036	13.15	3.581	24.32	24.23					
dc190240	12.36	3.408	24.62		Anova: Single Factor				
dc190243	12.36	3.385	24.46	24.54					
dc190253	13.05	3.645	24.94		<b>SUMMARY</b>				
dc190256	13.05	3.617	24.75	24.85	<i>Groups</i>	<i>Count</i>	<i>Sum</i>	<i>Average</i>	<i>Variance</i>
dc250237	12.74	3.456	24.22		Analyst 1	3	72.4465	24.14883	0.110458
dc250238	12.74	3.443	24.13	24.18	Analyst 2	3	73.56504	24.52168	0.111666
p2907017	11.81	3.228	24.41		Analyst 3	3	74.36278	24.78759	0.119366
p2907019	11.81	3.231	24.43	24.42					
p2907030	10.89	3.059	25.08						
p2907031	10.89	3.064	25.13	25.10					
p2907036	10.57	2.937	24.81		<b>ANOVA</b>				
p2907039	10.57	2.943	24.86	24.84	<i>Source of Variation</i>	<i>SS</i>	<i>df</i>	<i>MS</i>	<i>F</i>
					Between Groups	0.617737	2	0.308868	2.713424
					Within Groups	0.682979	6	0.11383	
					<i>P-value</i>	<i>F crit</i>			
					0.144769	5.143249			

**From ANOVA**

Repeatability standard deviation is the root of MS (mean square) Within Groups

$$\sqrt{0,11383} = 0,34 \% \text{ (w/w) or RSD } 1,38 \%$$

To calculate the within-lab reproducibility you first need to calculate the between group standard deviation  $s_x$  where n is the number of groups..

$$s_x = \sqrt{\frac{MKV_{\text{Betweengroups}} - MKV_{\text{Withingroups}}}{n}} = \sqrt{\frac{0.308869 - 0.11383}{3}} = 0.25\% \text{ (w/w)}$$

$$RSD_x = 1,04 \%$$

$$RSD_{Rw} = \sqrt{RSD_r^2 + RSD_x^2} = \sqrt{1.38^2 + 1.04^2} = 1.72\%$$

<b>Results:</b>	<b><i>RSD<sub>r</sub></i></b> =1.12% and from ANOVA 1.4 % <b><i>RSD<sub>Rw</sub></i></b> =1.72%
<b>Conclusions:</b>	Fit for purpose



Parameter	Description	
<b>Bias</b>	Laboratory bias Selectivity	
	<b>Demand</b>	
	Laboratory bias-<10% (relative) Selectivity -No co-elution found with methanol solvent, clomipramine internal standard or any of the compounds encountered frequently in cocaine street samples	
	<b>Experiment</b>	
	Laboratory bias- Cocaine certified reference sample run over a period of time and % bias determined. Selectivity -Methanol solvent, clomipramine internal standard and compounds such as lignocaine frequently in cocaine street samples were run using the procedure to check that co –elution did not occur.	
	<b>Evaluation</b>	
	Reference 99 % cocaine	
	Bias (Abs)	Bias (%)
	result	
	96.1446	-2.86
	99.0826	0.08
	105.3841	6.38
	101.8351	2.84
	94.5851	-4.41
	94.5042	-4.50
	99.2506	0.25
	94.8200	-4.18
	Avg=	-0.80
	SD =	3.95
	Relative bias is $0,80/99*100= -0.81\%$ (relative) with a standard deviation of 4.0 % (relative)	
	Selectivity - The results generated from running methanol solvent, clomipramine internal standard and compounds such as lignocaine were evaluated.	
<b>Results:</b>	Laboratory bias is - 0.81 % (relative) Selectivity -No co-elution found with methanol solvent, clomipramine internal standard or any of the compounds encountered frequently in cocaine street samples	
<b>Conclusions :</b>	Fit for purpose	

Parameter	Description																																						
Measuring range	Linearity Limit of Quantification (LOQ)																																						
	<b>Demand</b>																																						
	Linearity Residual maximum deviation <2% relative in the higher range – over 10 % cocaine																																						
	Limit of Quantification 1% with a signal to noise ratio >10																																						
	<b>Experiment</b>																																						
	Linearity Cocaine calibration standards set up and run (done in triplicate initially and augmented with further calibration curves generated for casework). When all standards run each is added into the calibration curve using the data analysis software.  % Residual = (predicted value for calibration standard- actual value) divided by actual value times 100. Actual value is the actual amount of calibration standard weighed out (and used as part of the calibration curve). Predicted value is the result obtained when this standard's result is recalculated using the calibration curve.																																						
	Comment on the procedure. Samples of the check standard (used in repeatability/reproducibility studies above) and the known certified reference material (as used for system bias) are run to ensure that the curve is accurate																																						
	Limit of Quantification Prepare a 1% w:w cocaine reference material : matrix mixture (Note the composition of the matrix will be determined by factors such as which constituents are normally present in street cocaine samples)  1% sample run a minimum of three times on GC-FID instrument to see if the response is reproducible and the S/N ratio is > 10																																						
	<b>Evaluation</b>																																						
	Linearity % Residual value calculated from results obtained. Less than 2 % relative for calibrants over 10 % (w/w) cocaine  Limit of Quantification- 1% w:w cocaine reference material : matrix mixture evaluated to ensure the response obtained was reproducible with S/N ratio > 10.																																						
	<table border="1"> <thead> <tr> <th rowspan="2"></th> <th colspan="3">Baseline</th> <th colspan="3">1 % (w/w) cocaine</th> <th rowspan="2">Ratio</th> </tr> <tr> <th>low</th> <th>high</th> <th>diff</th> <th>low</th> <th>High</th> <th>diff</th> </tr> </thead> <tbody> <tr> <td>DC03072</td> <td>16.662</td> <td>16.726</td> <td><b>0.064</b></td> <td>16.75</td> <td>60.5</td> <td><b>43.75</b></td> <td><b>683</b></td> </tr> <tr> <td>DC03074</td> <td>16.56</td> <td>16.665</td> <td><b>0.105</b></td> <td>16.6</td> <td>62.3</td> <td><b>45.7</b></td> <td><b>435</b></td> </tr> <tr> <td>DC03075</td> <td>16.638</td> <td>16.72</td> <td><b>0.082</b></td> <td>16.6</td> <td>61.5</td> <td><b>44.9</b></td> <td><b>547</b></td> </tr> </tbody> </table>		Baseline			1 % (w/w) cocaine			Ratio	low	high	diff	low	High	diff	DC03072	16.662	16.726	<b>0.064</b>	16.75	60.5	<b>43.75</b>	<b>683</b>	DC03074	16.56	16.665	<b>0.105</b>	16.6	62.3	<b>45.7</b>	<b>435</b>	DC03075	16.638	16.72	<b>0.082</b>	16.6	61.5	<b>44.9</b>	<b>547</b>
	Baseline			1 % (w/w) cocaine			Ratio																																
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<b>Results:</b>	<p>Linearity  For all calibration curves  Residual maximum deviation figure &lt;2% relative obtained</p> <p>Limit of Quantification  1% w:w cocaine reference material : matrix mixture gave a reproducible result and S/N ratio &gt; 10</p>
<b>Conclusions:</b>	Fit for purpose

Parameter	Description
<b>Measurement Uncertainty</b>	<p>COMMENT: <i>estimation of measurement uncertainty is outside the scope of this guide. However from validation data it may be possible to estimate uncertainty and here is presented one approach. Measurement Uncertainty is here calculated from the validation data according to ISO 11352 Water quality — Estimation of measurement uncertainty based on validation and quality control data. The approach is also given in Nordtest 537 <a href="http://www.nordtest.info">www.nordtest.info</a></i></p>
	<p>The data needed is  A Within-Lab reproducibility  B Bias measurements</p>
	<p>The standard uncertainty <math>u_c</math> can be calculated</p>
	$u_c = \sqrt{u(R_w)^2 + u(bias)^2}$
	<p>Where <math>u(R_w)</math> in this case is <math>RSD_{Rw}</math> and</p>
$u(bias) = \sqrt{bias^2 + \left(\frac{s_{bias}}{\sqrt{n}}\right)^2} + u(Cref)^2$	
<p>Where <math>u(Cref)</math> is the standard uncertainty for the reference value</p>	
<p>The expanded uncertainty (U) is obtained by multiplying the combined standard uncertainty, <math>u_c(y)</math>, by a chosen coverage factor, <math>k</math>. The expanded uncertainty provides an interval which may be expected to include a large proportion of the distribution of values which could be reasonably attributed to the measurand. For most purposes a value of 2 should be chosen for <math>k</math>.</p>	
$U = u_c(y) * 2$	
<b>Demand</b>	<p>Less than 20 % relative in the measurement range over 5 % (w/w) cocaine</p>
<b>Experiment</b>	<p><b>A: Within-laboratory reproducibility</b>  See precision section</p> <p><b>B. Bias</b>  See Bias section</p>
<b>Evaluation</b>	

	<p><b>A: intermediate precision.</b> From the precisions experiment <math>RSD_{Rw}=1.72\%</math></p> <p><b>B. Bias</b> The relative bias obtained with 9 measurement different days is -0,81% (relative) with a standard deviation of 4.0 % (relative)</p> <p>Purity assay: &gt;98.5% (w/w)</p> <p>The purity is taken to be <math>99.25 \pm 0.75\%</math> (w/w) with a rectangular distribution (<i>i.e. equal probability that the true value lies anywhere within the <math>\pm</math> range</i>).</p> <p>To obtain the relative standard uncertainty for purity, <math>u(ref)</math>, with a rectangular distribution the <math>\pm</math> value is divided by <math>\sqrt{3}</math>.  <math>u(ref) = 0.75/\sqrt{3}/99*100 = 0.44\%</math> (relative)</p> $u(bias) = \sqrt{bias^2 + \left(\frac{s_{bias}}{\sqrt{n}}\right)^2 + u(Cref)^2} = \sqrt{-0.81^2 + \left(\frac{4.0}{\sqrt{9}}\right)^2 + 0.44^2} = 1.6\%$ <p>=</p> <p><b>Calculation of the Combined &amp; Expanded Uncertainty</b>  <math>U = 2u_c = \sqrt{u(R_w)^2 + u(bias)^2} = \sqrt{1.72^2 + 1.6^2} = 4.7\%</math></p>
<b>Results:</b>	Expanded Uncertainty - Less than 15% relative in the measurement range over 5 %
<b>Conclusions:</b>	Fit for purpose

## 6. Conclusions

<b>Summary</b>	Laboratory/customer demands met	
	Results from casework calibration curves casework can be added to linearity studies	
<b>Plan of internal quality control:</b>	Control charts put in place to ensure the system is operating within its specifications. It involves regularly monitoring the result obtained of the concentration of the control check standard (used for repeatability and reproducibility studies above).	
<b>Statement on fitness for intended purpose</b>	<input checked="" type="checkbox"/> <b>The method fulfils the demand</b>	<input type="checkbox"/> <b>The method does not fulfil the demand</b> <b>The following should be done....</b>

### Reference Documents

EURACHEM / CITAC Guide CG4, “*Quantifying uncertainty in analytical measurement*”, second edition, 2000.  
 ENFSI QCC, “*Guidance for uncertainty measurement in quantitative analysis or testing (UM)*”, QCC-UM-001, Nov 2006.

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## 5.1b. Primarily Qualitative

### 5.1.4 Qualitative screening for gun shot residues

## IN-HOUSE VALIDATION OF MEASUREMENT PROCEDURES

# VALIDATION PLAN






### 1. Specify the measurement procedure, analyte, measurand and units

<i>The measurement procedure</i>	Gunshot Residue Analysis by Scanning Electron Microscopy / Energy Dispersive X-ray Spectrometry
<i>The measurand</i>	Number of PbSbBa particles with a diameter 0,5 $\mu$ m


### 2. Specify the Scope

<i>Matrix</i>	Surface of microscopic stub
<i>Measuring range</i>	0 ÷ 100 PbSbBa particles
<i>Intended use of the results</i>	Courts, prosecutors' and police intelligence reports

### 3. Requirement on the measurement procedure

<i>Parameters to be validated</i>		Value requested	
<i>Precision</i>	<i>Repeatability</i>		$RSD_r \leq 10 \%$
	<i>Within-lab reproducibility</i>		$RSD_{Rw} \leq 10 \%$
<i>Trueness</i>	<i>Test for bias</i>		$Bias \leq 15 \%$
	<i>Sensitivity</i>		$Sensitivity \geq 85\%$
<i>Measurement range</i>	<i>LOD</i>		$Particle's\ diameter \geq 0.5\ mm$

### 4. Origin of the Measurement Procedure

		<i>VALIDATION</i>
<i>New In-House Method</i>		<i>Full</i>
<i>Modified Validated/Standard Method</i>		<i>Partial, modified ASTM 1588 – 08 method</i>
<i>Official Standard Method</i>		<i>“Simple”</i>

## VALIDATION PLAN AND RESULTS

The whole process relies on results achieved from examination of ENSFI GSR PT test sample. In the example the test sample, coded as SPS-5P-1A, originates from GSR 2003 PT edition. It is the synthetic stub with known quantity and localization of PbSbBa particles on its surface. The test sample plays a role of a standard, however is treated as a regular sample.

Sample code	Total number of PbSbBa	Number of PbSbBa with a diameter of				
		10 □m	2.4 □m	1.2 □m	0.8 □m	0.5 □m
SPS-5P-1A	100	3 <sup>*</sup>	27 <sup>**</sup>	26 <sup>**</sup>	25 <sup>**</sup>	22 <sup>**</sup>
* only for sample orientation and focusing						
** assigned, known number of particles with given diameter						

Parameter	Description
<b>Precision</b>	Repeatability, Within Lab Reproducibility,
	<b>Demand</b>
	Repeatability $\leq 10\% RSD_r$ Within Lab Reproducibility $\leq 10\% RSD_{Rw}$
	<b>Experiment</b>
	<b>RSD<sub>r</sub></b> – The test sample was analysed automatically 6 times by the same operator and with the help of the same instrument. <b>RSD<sub>Rw</sub></b> – The test sample was examined once per month as the control sample. The same instrument was used, however various operators were involved.
	<b>Evaluation</b>



**1. RSD<sub>r</sub>**

Sample Name	Total number of PbSbBa	Number of PbSbBa with a diameter of			
		2.4 $\mu$ m	1.2 $\mu$ m	0.8 $\mu$ m	0.5 $\mu$ m
Run nr 1	92	26	26	24	16
Run nr 2	89	27	23	21	18
Run nr 3	89	26	24	23	16
Run nr 4	91	27	23	23	18
Run nr 5	88	26	25	22	15
Run nr 6	92	25	22	25	20
<b>AVG</b>	<b>90</b>	<b>26</b>	<b>24</b>	<b>23</b>	<b>17</b>
<b>SD</b>	<b>1,7</b>	<b>0,8</b>	<b>1,5</b>	<b>1,4</b>	<b>1,8</b>
<b>RSD<sub>r</sub></b>	<b>2%</b>	<b>3%</b>	<b>6%</b>	<b>6%</b>	<b>11%</b>

**2. RSD<sub>Rw</sub>**

Sample Nae	Total number of PbSbBa	Number of PbSbBa with a diameter of			
		2.4 $\mu$ m	1.2 $\mu$ m	0.8 $\mu$ m	0.5 $\mu$ m
January	87	25	25	22	15
February	86	27	25	20	14
March	87	27	24	19	17
April	94	27	25	24	18
May	86	26	24	21	15
June	89	27	22	24	16
July	87	26	24	20	17
August	2	27	26	22	17
September	90	26	23	22	19
October	90	26	26	21	17
November	88	25	25	22	16
December	86	27	25	20	14
<b>AVG</b>	<b>89</b>	<b>26</b>	<b>25</b>	<b>21</b>	<b>16</b>
<b>SD</b>	<b>2,6</b>	<b>0,8</b>	<b>1,2</b>	<b>1,6</b>	<b>1,5</b>
<b>RSD<sub>Rw</sub></b>	<b>3%</b>	<b>3%</b>	<b>5%</b>	<b>8%</b>	<b>9%</b>

**Conclusions: Fit for purpose**

<b>Parameter</b>	<b>Description</b>
<b>Trueness</b>	<p>SENSITIVITY – it is method's ability for identifying positive results. In the example, a positive result means correctly detected and classified PbSbBa particle. The sensitivity parameter is defined as</p> $\text{SENSITIVITY} = \frac{\text{TP}}{\text{TP} + \text{FN}} * 100\%$ <p>where TP – means number of true positive results, FN – means number of false negative results.</p> <p>Number of true positive TP results was calculated from the equation:</p> $\text{TP} = \text{Total Number of PbSbBa} - \text{FP}$ <p>The false positive result FP mostly may happen when the same particle is counted at least two times because of fields overlapping effect. False negative result FN encounters when PbSbBa particle is assigned to Pb, Ba, Sb, PbSb, PbBa, SbBa, Unclassified class or it stays undetected at all. Acquiring the true negative result TN is unlikely due to the fact that the test stub is non-PbSbBa particle free.</p> <p>The artificial ENFSI GSR test samples were used in ENFSI GSR PT since 1999 and could be used for validation and quality control.</p> <p>BIAS is defined as difference between lab's results and assigned value of particles on the test stub and it is expressed in the relative form.</p> $\text{BIAS} = \frac{\text{Total Number of PbSbBa} - \text{Assigned Number of PbSbBa}}{\text{Assigned Number of PbSbBa}} * 100\%$
	<b>Demand</b>
	<p>SENSITIVITY <math>\geq</math> 85%</p> <p>  BIAS   <math>\leq</math> 15%</p>
	<b>Experiment</b>
	Sensitivity and bias was calculated from data acquired from within lab reproducibility data.
	<b>Evaluation</b>

## 1. SENSITIVITY and BIAS

Sample Name	Method's performance among groups															
	2.4 □ m				1.2 □ m				0.8 □ m				0.5 □ m			
	FN	TP	Sensitivity [%]	BIAS [%]	FN	TP	Sensitivity [%]	BIAS [%]	FN	TP	Sensitivity [%]	BIAS [%]	FN	TP	Sensitivity [%]	BIAS [%]
January	2	25	93	-7	1	25	96	-4	3	22	88	-12	7	15	68	32
February	0	27	100	0	1	25	96	-4	5	20	80	-20	8	14	64	36
March	0	27	100	0	2	24	92	-8	6	19	76	-24	5	17	77	23
April	0	27	100	0	1	25	96	-4	1	24	96	-4	4	18	82	18
May	1	26	96	-4	2	24	92	-8	4	21	84	-16	7	15	68	32
June	0	27	100	0	4	22	85	-15	1	24	96	-4	6	16	73	27
July	1	26	96	-4	2	24	92	-8	5	20	80	-20	5	17	77	23
August	0	27	100	0	0	26	100	0	3	22	88	-12	5	17	77	23
September	1	26	96	-4	3	23	88	-12	3	22	88	-12	3	19	86	14
October	1	26	96	-4	0	26	100	0	4	21	84	-16	5	17	77	23
November	2	25	93	-7	1	25	96	-4	3	22	88	-12	6	16	73	27
December	0	27	100	0	1	25	96	-4	5	20	80	-20	8	14	64	36
		<b>AVG</b>	<b>98</b>	<b>-3</b>			<b>94</b>	<b>-6</b>			<b>86</b>	<b>-14</b>			<b>74</b>	<b>-26</b>
		<b>SD</b>	<b>3</b>	<b>3</b>			<b>4</b>	<b>4</b>			<b>6</b>	<b>6</b>			<b>7</b>	<b>7</b>


Sample Name	Total number of PbSbBa	Overall method's performance				
		FP	FN	TP	SENSITIVIT Y [%]	BIAS [%]
January	87	0	13	87	87	-13
February	86	0	14	86	86	-14
March	87	0	13	87	87	-13
April	94	0	6	94	94	-6
May	86	0	14	86	86	-14
June	89	0	11	89	89	-11
July	87	0	13	87	87	-13
August	92	0	8	92	92	-8
September	90	0	10	90	90	-10
October	90	0	10	90	90	-10
November	88	0	12	88	88	-12
December	86	0	14	86	86	-14
				<b>AVG</b>	<b>89</b>	<b>-12</b>
				<b>SD</b>	<b>3</b>	<b>3</b>

The overall bias for a typical samples is better than 15 %.

However for the smaller the fraction the higher bias and for the smallest fraction the bias is -26 %.

**Conclusions: Fit for purpose**

## 6. Conclusions

<b>Summary</b>	<p>This method is fit for the intended use since:</p> <ul style="list-style-type: none"> <li>▪ allows detecting particles of 0.5 <math>\mu</math>m in diameter.</li> <li>▪ repeatability and reproducibility expressed as <math>RSD_r</math> and <math>RSD_{RW}</math> is less than 10%.</li> <li>▪ sensitivity of the method is better than 85% and absolute value of bias is smaller than 15%.</li> </ul>	
<b>Plan of internal quality control:</b>	<p>Data collected during in within lab reproducibility experiment could be utilized for creating control chart where further results for the test sample can be regularly monitored.</p>	
<b>Statement on fitness for intended purpose</b>	 <p><b>The method fulfills the demand</b></p>	<p><input type="checkbox"/> <b>The method does not fulfill the demand</b>  <b>The following should be done....</b></p>

## 5.1.5 DNA-kit 17-loci STR PCR chemistry

### SINGLE LABORATORY VALIDATION OF MEASUREMENT PROCEDURES (Amplification System -NGMSElect)

*Sample validation-Guideline purposes only*

#### 1. Specify the measurement procedure, analyte, measurand and units

<i>The measurement procedure</i>	Determine the ability of NGM SElect (Life Technologies) STR procedures to generate DNA profiles from a range of sample types.
<i>Analyte</i>	Human Genomic DNA
<i>The measurand</i>	17-loci STR DNA Profile
<i>Unit</i>	N/A

#### 2. Specify the Scope

<i>Matrix</i>	Crimestain and reference DNA samples
<i>Measuring range</i>	~125pg/ul-1ng/ul

#### 3. Requirement on the measurement procedure

<i>Intended use of the results</i>	<p>The result of using NGM SElect chemistry is that more informative (i.e. containing more loci and therefore allele calls) DNA profiles should be generated from samples than is currently possible with SGM+ profiles.</p> <p>The overall goal is to set up DNA profiling reactions using improved chemistry due to developments in human identity kit production, evidenced by quiet baselines and additional allelic information.</p>
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**Internal validation:** To ensure kit performs as expected in the laboratory.

<i>Parameters to be validated</i>	<i>Value requested</i>
Known and non-probative evidence samples (Concordance)	To confirm the SGM+ loci data generated using NGM SElect chemistry is equivalent to the profile data generated for the same samples using SGM+ chemistry.
Reproducibility and Repeatability	Confirm that the STR profiles generated using NGM SElect chemistry at 500 pg of 3 heterozygote DNA amplified at 30 cycles and 1 ng amplified at 29 cycles by 2 operators and analysed on the 3130 analyser are complete and accurate using GMIDX software.

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Precision of sizing (Match Criteria)	All sized and called alleles in the samples analysed to display a difference of less than 0.5 bp when compared to the size of the corresponding allele in the allelic ladder, with the largest size difference in any of the runs being 0.27 bp.
Sensitivity - Corroborate recommended input DNA concentration	Complete profiles generated from 125 pg input template DNA using both 29 and 30 cycle amplifications.
Stochastic Determination of allele call thresholds on GMIDX analysis software.	Confirm the range of template DNA amounts where allele dropout is observed. Set the appropriate allele calling thresholds on GMIDX analysis software.
Determination of stutter ratios/thresholds	Confirm the stutter ratios/thresholds down and up, for each locus with regard to the manufacturer's expected ratios from DNA in solution and DNA on FTA cards.
Peak balance analysis	Determine data for intra-locus and inter-locus peak height balances. Average intra-locus and inter-loci balance is greater than 60%.
Mixture Analysis	To determine that NGMSelect chemistry is more sensitive at detecting mixtures than SGM+ irrespective of the choice of 29 or 30 cycle amplification conditions.
Anticontamination	All profiles generated will be assessed for 1) the potential of drop in 2) any spurious peaks in the negative samples.
Qualifying test studies	The qualifying test will consist of at least 50 typical crime stain sample types to be amplified using NGM Select.

#### 4. Origin of the Measurement Procedure

		VALIDATION
<i>New In-House Method</i>	<b>X</b>	<i>Full internal validation</i>
<i>Modified Validated Method</i>	<input type="checkbox"/>	<i>Partial</i>
<i>Official Standard Method</i>	<input type="checkbox"/>	<i>"Simple"</i>



## 5. Performance Parameters

Parameter	Description
Known and non-probative evidence samples (Concordance)	To confirm the SGM+ loci data generated using NGM Select chemistry is equivalent to the profile data generated for the same samples using SGM+ chemistry.
	<b>Demand</b>
	The SGM+ loci data generated using NGM Select are 100% concordant with profile data generated for the same samples using SGM+ chemistry.
	<b>Experiment</b>
	50 population database samples (buccal swab samples on FTA) amplified using NGM Select, at 1ng at 29 cycles and analysed on the 3130 analyser.
	<b>Evaluation</b>
	Examine the profiles generated for concordance
<b>Results:</b>	The SGM+ loci data generated using NGM Select are 100% concordant with profile data generated for the same samples using SGM+ chemistry.
<b>Conclusions:</b>	Fit for purpose

Parameter	Description
Reproducibility and Repeatability	Confirm that the STR profiles generated using NGM Select chemistry at 500 pg of 3 heterozygote DNA amplified at 30 cycles and 1 ng amplified at 29 cycles by 2 operators and analysed on the 3130 analyser are complete and accurate using GMIDX software.
	<b>Demand</b>
	Complete and accurate (concordant) profiles generated
	<b>Experiment</b>
	500 pg of 3 heterozygote DNA samples amplified at 30 cycles and 1 ng amplified at 29 cycles by 2 operators and analysed on the 3130 analyser
	<b>Evaluation</b>
	Profiles generated analysed



<b>Results:</b>	All profiles generated from the three heterozygotic samples were concordant between repeats at 29 and 30 cycles and between operators
<b>Conclusions:</b>	Fit for purpose

Parameter	Description
Precision of sizing (Match Criteria)	Precision of sizing
	<b>Demand</b>
	All sized and called alleles in the samples analysed to display a difference of less than 0.5 bp when compared to the size of the corresponding allele in the allelic ladder, with the largest size difference in any of the runs being 0.27 bp.
	<b>Experiment</b>
	Resolution of the allelic ladder calculated across one injection of 12 ladder samples injected twice and the ladders used in the concordance, repeatability and reproducibility studies. The size range for each allele in the samples across the repeatability and reproducibility studies were calculated
	<b>Evaluation</b>
	Data compiled and analysed
<b>Results:</b>	<p>All sized and called alleles in the samples analysed displayed a difference of less than 0.5 bp when compared to the size of the corresponding allele in the allelic ladder, with the largest size difference in any of the runs being 0.27 bp.</p> <p>NGM SElect is precise, that is all alleles fall within a 0.5 bp window around the measured size for the corresponding allele in the allelic ladder.</p> <p>The 29 allelic ladders analysed all were able to resolve single base pair size differences. NGM SElect allelic ladder can resolve single base pair differences.</p>
<b>Conclusions:</b>	Fit for purpose

Parameter	Description
Sensitivity - Corroborate recommended input DNA concentration	Sensitivity - Determine the lowest DNA input amt per reaction that can generate a full 17 loci STR profile using NGM Select.
	<b>Demand</b>
	Complete profiles generated from 125 pg input template DNA using both 29 and 30 cycle amplifications.
	<b>Experiment</b>
	A 7 dilution series from 1ng to 10 pg amplified from three heterozygote samples on four separate reactions at 500 pg at 30 cycles and 1 ng at 29 cycles and analysed on the 3130 analyser. (84 samples for each input template and cycle number)
	<b>Evaluation</b>
	Data compiled and analysed
<b>Results:</b>	Complete profiles were generated from 125 pg input template DNA using both 29 and 30 cycle amplifications. Allele calls were also generated from template amounts of less than 125 pg using both 29 and 30 cycles. 30 cycle amplifications were able to generate more allele calls from low template amounts (50, 30 and 10 pg) than 29 cycle amplifications.
<b>Conclusions:</b>	Fit for purpose

Parameter	Description
Stochastic Determination of allele call thresholds on GMIDX analysis software.	Stochastic Determination of allele call thresholds
	<b>Demand</b>
	Confirm the range of template DNA amounts where allele dropout is observed. Set the appropriate allele calling thresholds on GMIDX analysis software.
	<b>Experiment</b>
	Samples generated during sensitivity studies used (i.e. dilution series)
	<b>Evaluation</b>
	Data from the dilution series in the sensitivity study examined based on the expected full STR profile.
<b>Results:</b>	Analysis of the data following injection on the 3130 analyser indicates the following allele calling thresholds when analysing 29 cycle NGM SE amplifications: Heterozygote calling threshold: 50 rfu Homozygote calling threshold: 270 rfu

<b>Conclusions:</b>	Fit for purpose

Parameter	Description
Determination of stutter ratios/ thresholds	Determination of stutter ratios/ thresholds
	<b>Demand</b>
	Confirm the stutter ratios/thresholds down and up, for each locus with regard to the manufacturer's expected ratio's from DNA in solution and DNA on FTA cards
	<b>Experiment</b>
	The 50 buccal FTA concordant samples were amplified with NGM SElect at 1ng at 29 cycles and analysed on the 3130 analyser with the calling threshold set at 1 rfu.
	<b>Evaluation</b>
	Data analysed
<b>Results:</b>	The stutter levels of the NGM SElect chemistry are in agreement with the manufacturers expected stutter levels. The laboratory will adopt the manufacturers recommendations for stutter levels and apply these values to the analysis of NGMSElect profiles until such time as enough data has been collected by the laboratory to change the values as appropriate (i.e. longitudinal study).
<b>Conclusions:</b>	Fit for purpose

Parameter	Description
Peak balance analysis	Determine data for intra-locus and inter-locus peak height balances
	<b>Demand</b>
	Average intra-locus and inter-loci balance is greater than 60%.
	<b>Experiment</b>
	Data from samples from the concordance, repeatability, reproducibility and sensitivity studies analysed with regard intra-locus and inter-locus peak height balances.
	<b>Evaluation</b>
	Data analysed
<b>Results:</b>	Average intra-locus and inter-loci balance found to be greater than 60%. Operators should be aware that instances where intra-locus balance values of less than 60% are observed are uncommon but not rare occurring approx 3% of the time.
<b>Conclusions:</b>	Fit for purpose

Parameter	Description
Mixture Analysis	Analysis of mixture samples. The highest mixture ratio that can generate detectable alleles for the minor contributor using both SGM+ and NGM SElect.
	<b>Demand</b>
	NGMSElect chemistry is more sensitive at detecting mixtures than SGM+ irrespective of the choice of 29 or 30 cycle amplification conditions.
	<b>Experiment</b>
	Ratios of 1:1 to 1:10 were amplified for two of the three heterozygote samples using both SGM+ and NGM SElect. These were amplified at 500 pg at 30 cycles and 1 ng at 29 cycles for the NGM SElect kit and analysed using the 3130.
	<b>Evaluation</b>
	Analyse data
<b>Results:</b>	NGMSElect chemistry is more sensitive at detecting mixtures than SGM+ irrespective of the choice of 29 or 30 cycle amplification conditions.
<b>Conclusions:</b>	Fit for purpose

Parameter	Description
Anticontamination	Assess contamination risk
	<b>Demand</b>
	All profiles generated will be assessed for 1) the potential of drop in 2) any spurious peaks in the negative samples
	<b>Experiment</b>
	All profiles generated assessed for 1) the potential of drop in 2) any spurious peaks in the negative samples
	<b>Evaluation</b>
	Data analysed
<b>Results:</b>	Allelic drop-in can occur at a low level with NGM Select chemistry. The samples observed had a low template input (100 pg and 50 pg) and were both amplified at 30 cycles. Allelic drop in is a reported phenomenon in the literature (Weiler et al, FSI Genetics, 2012, Jan) and care must be taken when interpreting profiles that contain alleles generated from low template inputs.
<b>Conclusions:</b>	Fit for purpose

Parameter	Description
Qualifying test studies	The qualifying test will consist of at least 30 typical crime stain sample types to be amplified using NGM SElect.
	<b>Demand</b>
	Good success rate (i.e. full DNA profiles obtained) and profiles give concordant results
	<b>Experiment</b>
	Typical casework type samples were selected for profiling with NGM SElect. These included cigarette butts, chewing gum, dilutions of semen on swabs, blood spots on denim fabric, Epithelial cells on swabs and minitapes, tissue samples and blood samples submitted for paternity testing.
	Reference samples and crimestains from CTS Forensic Biology DNA mixture trials were profiled with NGMSElect. The testing body only accepts the SGM+ loci which overlap with the NGMSElect and therefore only the typing information for these loci were submitted.  Samples were processed using the Qiagen EZ1XL machine
	<b>Evaluation</b>
	Data analysed
<b>Results:</b>	Expected profiles obtained from crimestain samples and CTS proficiency trials
<b>Conclusions:</b>	Fit for purpose

## 6. Conclusions

<b>Summary</b>	All parameters tested yielded fit for purpose results. Determination of stutter ratios/ thresholds data to be reviewed following longitudinal studies	
<b>Plan of internal quality control:</b>	Extraction positive and negative controls and amplification positive and negative controls in system  Appropriate level of anticontamination measures and environmental monitoring put in place	
<b>Statement on fitness for intended purpose</b>	<input checked="" type="checkbox"/> <b>The method fulfils the demand</b>	<input type="checkbox"/> <b>The method does not fulfil the demand</b> <b>The following should be done....</b>



**5.1.6 human salivary alfa-amylase detection by RSID™-saliva test.****IN-HOUSE VALIDATION OF  
MEASUREMENT PROCEDURES**

By courtesy by Crab R. and the DNA-unit of the NICC






**VALIDATION PLAN****1. Specify the measurement procedure, analyte, measurand and units**

<i>The measurement procedure</i>	Detection of human saliva using the RSID™-Saliva lateral flow immunochromatographic strip test.
<i>Analyte</i>	Human salivary alfa-amylase
<i>The measurand</i>	Qualitative detection of human saliva.
<i>Unit</i>	N/A





**2. Specify the Scope**

<i>Matrix</i>	Fluids, crime scene samples
<i>Measuring range</i>	
<i>Intended use of the results</i>	Detection / confirmation of human origin (saliva) at crime scene


**3. Requirement on the measurement procedure**

<i>Parameters to be validated</i>		<i>Value requested</i>
<i>Precision</i>	<i>Repeatability</i>	 Identical results
	<i>Within-lab reproducibility</i>	 Identical results
<i>Trueness</i>	<i>Specificity</i>	 Investigate the cross-reactivity with <ul style="list-style-type: none"> <li>• Saliva from animals</li> <li>• Other human body fluids</li> </ul>
<i>Measurement range<sup>2</sup></i>	<i>LOD</i> <i>High dose hook effect (upper LOD)</i>	 <1 µL <sup>2</sup>  determination

<sup>2</sup> The kit proposes 1 µL as LOD, but in practice it will be lower

<i>Ruggedness (Robustness)</i>	<i>Cotton swabs</i>		No influence
	<i>Forensic-like surfaces</i>		No influence
	<i>Stability of processed teststrips</i>	 	Readability after storage
STR analysis starting from the RSIDTM extraction buffer.	Genetic profile stability		No influence on the genetic profile using the extraction buffer

#### 4. Origin of the Measurement Procedure

		VALIDATION
<i>New In-House Method</i>		<i>Full</i>
<i>Modified Validated/Standard Method</i>		<i>Partial</i>
<i>Official Standard Method</i>		<i>“Simple”</i>

Parameter	Description				
Precision	Repeatability, Within.Lab Reproducibility,				
	<b>Demand</b>				
	Identical results				
	<b>Experiment</b>				
	Repeatability : A dilution series of liquid saliva diluted in RSID™ extraction buffer, processed by one person in triplicate.				
	Within Lab reproducibility : an identical dilution series as above, but by another technician and on another.				
	<b>DATA</b>				
		<b>Technician 1</b>	<b>Technician 2</b>		
	<b>Saliva concentration nL/µl</b>	<b>Run 1</b>	<b>Run 2</b>	<b>Run3</b>	<b>Run 4</b>
	<b>0.000</b>	-	-	-	-
<b>0.005</b>	-	-	-	-	
<b>0.024</b>	-	-	-	-	
<b>0.048</b>	-	-	-	-	
<b>0.200</b>	+	+	+	+	
<b>0.333</b>	+	+	+	+	
<b>0.500</b>	+	+	+	+	
<b>2.494</b>	+	+	+	+	
<b>4.975</b>	+	+	+	+	
<b>Results:</b>	Corresponding results				
<b>Conclusion</b>	Fit for purpose				

Parameter	Description
<b>Trueness</b>	Specificity
	<b>Demand</b>
	Investigate the cross-reactivity using the kit on saliva samples of animals.
	Investigate the cross-reactivity using the kit on other human body fluids
	<b>Experiment</b>
<p>A. Saliva samples of a wide variety of animals was tested (saliva collected from animals of friends and colleagues and obtained from animals in the Antwerp Zoo)</p> <p>B. Human body fluids tested:  Blood  Urine  Sperm  Mother milk  Nasal secretions</p> <p>The human body fluids were tested:  ○ separately  ○ in mixture with human saliva</p>	

**DATA**

A. all animal samples gave a negative test result except the saliva sample of the Gorilla (*Gorilla*).

B. Test results on other body fluids (separately or mixed)

Saliva (µL)	Blood (µL)	Urine (µL)	Sperm (µL)	Breast Milk (µL)	Result
	10				-
		10			-
			10		-
				10	(weak) +
	10	10	10		-
5	5				+
5	50				+
5		5			+
5		50			+
5			5		+
5			50		+

Addition tests following the cross-reactivity seen on breast milk.

Saliva (µL)	Blood (µL)	Urine (µL)	Sperm (µL)	Breast Milk (µL)	Result
				1	-
				5	-
				10	(weak)+
				20	+
				50	+
10				20	+
	10			20	(weak)+
		10		20	+
			10	20	+
	10	10	10	5	-
	10	10	10	50	+
10	10	10	10	20	+

Nasal secrete (which were collected with a cotton swab and not calculated in µL)

Nasal secrete	result
Swab 1	+
Swab 2	+

**Results:**

Cross-reactivity could be seen on the saliva of Gorilla. Cross-reactivity of the kit was demonstrated for breast milk and nasal secretes.

**Conclusions :**

The purpose of the kit is to detect traces on the crime scene which could be used for DNA-analysis. In this manner the cross-reactivity with Gorilla, breast milk and nasal secretes are not interfering the final purpose of the kit, only false positive for the validation of the kit in an analytical way. The false-positivity of nasal secretes is in routine use an advantage, because this samples can also be used for DNA-analysis.

Fit for purpose

Parameter	Description																													
Measurement range	Limit of detection																													
	High dose hook effect (upper limit of detection)																													
	<b>Demand</b>																													
	Determine the minimal amount of saliva needed to obtain a positive test.																													
	Determine the maximum amount of saliva still usable for this kit.																													
	<b>Experiment</b>																													
	<ul style="list-style-type: none"> <li>• Dilution series of human saliva in RSID™ saliva extraction buffer</li> <li>• Dilution series of human saliva in RSID™ saliva extraction buffer, but saliva from another testperson, because concentration of salivary alfa-amylase can differ between persons)</li>   <li>• Mixtures of undiluted saliva and RSID™ saliva extraction buffer.</li> </ul>																													
	<b>DATA</b>																													
	<b>LOD</b>																													
	<table border="1" style="width: 100%; border-collapse: collapse;"> <thead> <tr> <th style="width: 33%;">Salivaconcentration nL/µl</th> <th style="width: 33%;">Saliva from testperson1</th> <th style="width: 33%;">Saliva from testperson2</th> </tr> </thead> <tbody> <tr><td>0.000</td><td>-</td><td>-</td></tr> <tr><td>0.005</td><td>-</td><td>-</td></tr> <tr><td>0.024</td><td>-</td><td>-</td></tr> <tr><td>0.048</td><td>-</td><td>weak +</td></tr> <tr><td>0.200</td><td>+</td><td>+</td></tr> <tr><td>0.333</td><td>+</td><td>+</td></tr> <tr><td>0.500</td><td>+</td><td>+</td></tr> <tr><td>2.494</td><td>+</td><td>+</td></tr> <tr><td>4.975</td><td>+</td><td>+</td></tr> </tbody> </table>	Salivaconcentration nL/µl	Saliva from testperson1	Saliva from testperson2	0.000	-	-	0.005	-	-	0.024	-	-	0.048	-	weak +	0.200	+	+	0.333	+	+	0.500	+	+	2.494	+	+	4.975	+
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Salivaconcentration (nL/µl)	Saliva from testperson1	Saliva from testperson2																												
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48	+	+																												
91	+	+																												
200	+	+																												
333	+	+																												
500	+	+																												
<b>Results:</b>	<p>The limit of detection of the testkit is a saliva concentration of 0.2 nL/µL extraction buffer. Recalculation results in a corresponding value of 0.004 µL saliva addition to the teststrip.</p> <p>The high dose hook effect could not be seen at high concentration (500 nL/µl). recalculation results in a corresponding value of 10 µL saliva addition directly (without the dilution step) to the teststrip. (2500x LOD)</p>																													
<b>Conclusions:</b>	Fit for purpose																													

Parameter	Description
<b>Ruggedness</b>	Ruggedness
	<b>Demand</b>
	What is the influence on results of the RSID™ saliva kit by : <ol style="list-style-type: none"> <li>1. the oral swab used as collector of the trace</li> <li>2. the surface/object where the trace is collected from</li> <li>3. storage of processes test strips</li> </ol>
	<b>Experiment</b>
	<ol style="list-style-type: none"> <li>1. A dilution series of liquid saliva was deposite on sterile cotton swabs and allowed to air-dry prior to processing.</li> <li>2. A range of routinely encountered forensic exhibits were laboratory prepared (normal use like drinking, smoking, wetting and wearing) by different testpersons (called G, M and J)               <ul style="list-style-type: none"> <li>○ Aluminium drinking cans</li> <li>○ Cigarette butts</li> <li>○ Envelopes</li> <li>○ Balaclava</li> </ul> <p>A range of clothes, positively prepared in the laboratory by depositing. Clothes were laboratory control prepared with difference concentration of saliva and air-dryed. Those clothes were screened with the Phadebas Forensic Press test, and possible traces were collected</p> </li> <li>3. A collection (n=27) of RSID saliva tests was processed and stored afterwards for one year at a dark, dry place at room temperature. The reading of the strips was compared to the initially obtained results.</li> </ol>



**DATA**

1. The addition of diluted saliva on the cotton swab followed by air-drying and subsequently processed by extraction using the RSID™ extraction buffer resulted in a higher LOD for the kit. Whereas the LOD for liquid saliva was the deposit of the equivalent of 0.004 µL saliva on the test strip, the equivalent of saliva to be added starting from air-dried swabs 0.3µL.

2.

Item	Test person	Result	Test person	Result	Test person	Result
Aluminum soda can	G	+	M	+	J	+
	G	+	M	+		
Cigarette butt (1x1 cm)	G	+				
	G	+				
Cigarette butt (3x3 mm)	G	+				
	G	+				
Envelope (1x1 cm)	G	+	M	+	J	+
Envelope (3x3 mm)	G	+	M	+	J	+
Balaclava-mouth (3x3 mm)	G	weak+	M	weak+	J	+
Balaclava-nose (3x3 mm)	G	weak+	M	weak+	J	weak+

Different volumes of saliva were added to cotton T-shirt, a woolen jumper and a jeans, followed by air-drying. The Phadebas Forensic Press test was used to localize possible saliva. The localized places were extracted by cutting out a 2x2 mm trace, which was processed.

In general can be shown that traces (by depositing of 0.5 µL saliva) could be confirmed by the RSDTM saliva test. There is however a tendency that the detection of possible traces differs using different types of textiles, however this is depending on the efficiency of the Phadebas Forensic Press test, which is outside the scope of this validation study.

3. All strips were still presenting the same result as the initial reading after a storage of one year.

**Results:**

The difference between the use of liquid saliva or a cotton swab used during sampling is resulting in a higher LOD of the test, however still with a LOD low enough for routine use (0.3 µL).

Different types of forensic-like samples were tested and the RSID™ saliva kit is able to detect saliva from samples likely to be encountered in forensic laboratory case work.

If a rereading is necessary the storage of the strips at a dark, dry place at room temperature is possible to obtain stable results.


**Conclusions:**

Fit for purpose



Parameter	Description
<b>Suitability in the following DNA-analysis</b>	Investigate the suitability of the RSID™ kit prior to DNA-STR analysis
	<b>Demand</b>
	No influence of the RSID™ extraction buffer on the resulting genetic profile.
	<b>Experiment</b>
	<p>All samples created during this validation study were investigated by DNA-STR analysis.</p> <p>Samples resulting in a positive RSID™ saliva test result were resulting in a full genetic profile.  Samples with values lower the LOD (a very weak or negative test result, 0.048 nL/μl dilution as an example) were resulting to a partial or negative genetic profile.</p> <p>The study of the surfaces used for forensic analysis used two different sampling sizes of cigarette butts, envelopes and textiles and both were resulting to full genetic profiles.</p>
<b>Results:</b>	If a positive signal is obtained by the RSID™ saliva kit, a full genetic profile can be obtained.
<b>Conclusions :</b>	Fit for purpose

## 6. Conclusions

<b>Summary</b>	Laboratory/customer demands met	
<b>Plan of internal quality control:</b>	Each new serial number of kits will be tested using a positive control (10 µL liquid saliva added to 200µL RSID™ extraction buffer) and a negative control (only RSID™ extraction buffer).	
<b>Statement on fitness for intended purpose</b>	 <p>The method fulfills the demand</p>	<input type="checkbox"/> <b>The method does not fulfill the demand</b> <b>The following should be done....</b>

## Reference Documents

## 5.2. Human-based examples

The two examples are provided with the aim to demonstrate Demonstration of how the generic criteria are have been applied to two specific forensic evaluation fields: the fingermark and fingerprint comparison and the bullet and cartridge cases comparison. The fingerprint field is the example of a newly developed when the firearms field is an already long established field. These examples have to be considered as two specific solutions adapted particular situations and not as the solution to be followed step by step independently of the situation and environment.

In order to achieve validation, both fields of expertise have followed a similar approach. The first step consisted in a review of the scientific literature, in order to compile the existing and established knowledge within the respective fields. A short list of references has been extracted to support the different aspects of the method development. In house research and development projects have been run to cover the aspects not existing in the literature and the results have been published. The performance of the practitioners is monitored along the ACEV examination through specific solutions in the protocol, for example to address the questions of specimen usability, repeatability, reproducibility and context bias management. Proficiency tests are used to test the global performance of the practitioners. Finally an education program is developed to ensure, maintain and check the competence of the practitioners.

### 5.2.1 Fingermark and fingerprint comparison

This example is derived from the new service (2012)“probabilistic fingerprint interpretation” of the fingerprint individualization group of the Netherlands Forensic Institute.

#### 5.2.1.1. Method development

Dactyloscopy is on one of the oldest fields in forensic science. The description, the classification and the selection of the features are abundantly mentioned in the literature of the XIX<sup>th</sup>, XX<sup>th</sup> and XXI<sup>th</sup> century.

#### Feature description

Ashbaugh, D.R., *Qualitative-Quantitative Friction Ridge Analysis – An Introduction to Basic and Advanced Ridgeology*. Practical Aspects in Criminal and Forensic Investigations, ed. V.J. Geberth. 1999, Boca Raton: CRC Press.

#### Feature selection and classification

Hutchins, L.A., *Systems of friction ridge classification*, in *The Fingerprint Sourcebook*, E.H. Holder, L.O. Robinson, and J.H. Laub, Editors. 2011: Washigton D.C., USA.  
United States Department of Justice and Federal Bureau of Investigation, *The Science of Fingerprints*. 1984, Washington DC: U.S. Government Printing Office.

#### Feature availability, measurability and specimens usability

Langenburg, G.M., *A Critical Analysis and Study of the ACE-V Process*, in *School of Forensic Science*, 2012, University of Lausanne, Switzerland, 48HrBooks, USA. p. 355.

#### Feature distinctiveness

The distinctiveness of the fingermark and especially of the minutiae configurations (2<sup>nd</sup> level of detail) is described in the literature. But research is still ongoing on how to compute a more objective strength of evidence for the 1<sup>st</sup>, 2<sup>nd</sup> and 3<sup>rd</sup> levels of detail, and about the dependence between the different levels of details.

#### General

Stoney, D.A., *Measurement of Fingerprint Individuality*, in *Advances in Fingerprint Technology*, H.C. Lee and R.E. Gaensslen, Editors. 2001, CRC Press: Boca Raton. p. 327-387.

Neumann, C., *Statistics and Probabilities as a Means to Support Fingerprint Examination*, in *Lee and Gaensslen's Advances in Fingerprint Technology*, R. Ramotowski, Editor. 2012, CRC Press. p. 407 - 452.

Langenburg, G., *Scientific Research Supporting The Foundations of Friction Ridge Examinations*, in *The Fingerprint Sourcebook*, E.H. Holder, L.O. Robinson, and J.H. Laub, Editors. 2011, U.S. Department of Justice, NIJ: Washington D.C., USA. p. 14-3 – 14-31.

#### **1<sup>st</sup> level details**

Haraksim, R., et al. *Assignment of the evidential value of a fingermark general pattern using a Bayesian network*. In: *International Conference of the Biometrics Special Interest Group - (BIOSIG)*. 2013. Darmstadt, Germany: Gesellschaft für Informatik e. V. (G.I.).

de Jongh, A., et al., *The frequencies of arch, loop and whorl fingerprint patterns. An empirical study in the Dutch population*. NFI Internal report / draft for publication, 2013.

#### **2<sup>nd</sup> level details**

Gutiérrez-Redomero, E., et al., *Are the population differences in minutiae frequencies? A comparative study of two Argentinian population samples and on Spanish sample*. *Forens. Sci. Int.*, 2012. **222**: p. 266-276.

#### **3<sup>rd</sup> level details**

Anthonioz, A., et al., *Level 3 Details and Their Role in Fingerprint Identification: A Survey among Practitioners*. *J. Forens. Ident.*, 2008. **58**(5): p. 562 - 589.

#### **Feature dependence**

Further research is necessary to understand the degree of dependence of the information of each of the 3 levels of details in order to combine them in a logically correct manner.

#### **Feature permanence**

Barns, J., *History*, in *The Fingerprint Sourcebook*, E.H. Holder, L.O. Robinson, and J.H. Laub, Editors. 2011, U.S. Department of Justice, NIJ: Washington D.C., USA. p. 1-5 – 1-22.

#### **Feature robustness**

Champod, C., et al., *The Friction Ridge Identification Process*, in: *Fingerprints and other Ridge Skin Impressions*, C. Press, Editor. 2004, CRC press: London.

#### **5.2.1.2. Validation of the performance**

##### **Practitioner reproducibility**

The reproducibility is addressed in each case, 2 examiners encode the specimens (marks and prints) separately and then merge their results. The rules are in Quality document: “the “description of the fingermark comparison method””.

Haraksim, R. and D. Meuwly, *Fingerprint examiners: a comparison system whose performance can be measured*. NFI Internal report / draft for publication, 2013.

##### **Practitioner repeatability**

Langenburg, G. and C. Champod, *The GYRO System - A recommended Approach to More Transparent Documentation*. *Journal of Forensic Identification*, 2010. **61**(4): p. 373 - 384.

##### **Specimen usability**

The criteria to determine the suitability for comparison is not made explicit yet, but it will be added to the quality document “description of the fingermark comparison method”. For every individual case the usability is documented in the case notes.

Ulery, B.T., et al., *Understanding the sufficiency of information for latent fingerprint value determinations*. *Forens. Sci. Int.*, 2013. **230**(1-3): p. 99-106.

#### **Proficiency test**

The fingerprint individualization group takes part twice a year the CTS proficiency test and once a year to the ENFSI-EFPWG collaborative exercise.

## Context bias management


The management of the case information and the sequential unmasking are described in the draft quality document: “Comparative dactyloscopic examination”.

### 5.2.1.3. Assessment of the competence of the practitioner

The competence of the practitioner is assessed during a training and assessment programme. Passing the assessment leads to the authority to sign reports. The training programme is composed of general courses (criminalistics, forensic statistics, law, report writing and courtroom training) as well as area specific training on the job and other training activities (literature reading, conference attendance).

The assessment of the competence is performed through written and oral examinations, assessment of the practical work and the defence of casework reports in front of a moot court constituted of professionals (judge, lawyer, counter expert). The final stage is constituted by the defence of casework reports in front of an examination commission composed of an external expert of the relevant area, a prosecutor and an internal criminalistics expert. Every four years the signed-off forensic practitioners are reassessed defending a new set of reports in front a similar examination commission.

The general competences that are relevant to all fields of expertise are described in the “Education manual – training and assessment program for the certification forensic practitioners”. The area specific competences relevant to the field of expertise of the candidate are detailed in a field specific and personalized training and assessment program. Such a program is composed of training activities composed of simulated casework using portfolios of known cases, real casework, collaborative exercises and proficiency tests. During the training the supervision decreases and the independency of the candidate increases. The tutor of the candidate monitors the results of the training activities

<b>Summary</b>	Laboratory/customer demands met	
<b>Statement on fitness for intended purpose</b>	 <b>The method fulfills the demand</b>	<input type="checkbox"/> <b>The method does not fulfill the demand</b> <b>The following should be done....</b>

## 5.2.2 Bullet and cartridge case comparison

This example is derived from the service “Comparison of spend ammunition part with or without a firearm” developed from the 1950s with further development on the methodology from 2008 by the firearms and ammunition group of the Netherlands Forensic Institute.

### 5.2.2.1. Method development

The bullet and cartridge cases comparison is one of the traditional fields of forensic examination for which the description, the classification and the selection of the features abundantly described in the literature.

#### Feature description, selection, classification

Rowe, W.F., *Firearms laboratory analysis*, in *Encyclopedia of Forensic Sciences*, J.A. Siegel, P.J. Saukko, and G.C. Knupfer, Editors. 2000, Academic Press: London. p. 944-949.

Dutton, G., *Firearms: Bullet and Cartridge Case Identification*. Wiley Encyclopedia of Forensic Science, 2009.

Rivera, G.C., *Subclass Characteristics in Smith & Wesson SW40VE Sigma Pistols*. AFTE Journal, 2007. **39**(3): p. 253-258.

Saribey, A. and A. Grace Hannam, *Comparison of the Class and Individual Characteristics of Turkish 7.65 mm Browning/.32 Automatic Caliber Self-Loading Pistols with Consecutive Serial Numbers*. . Journal of Forensic Sciences 2013. **58**(1): p. 146-150.

#### Feature availability, measurability and specimens usability

Kerstholt, J.A., et al., *Does suggestive information cause a confirmation bias in bullet comparisons?* Forensic Science International, 2010(198): p. 138–142.

#### Feature distinctiveness

The distinctiveness of the features is discussed in the literature, but more research is necessary to compute a more objective strength of evidence for the different features present on the bullets and cartridge cases and about the dependence between these features.

Mikko, D., J. Miller, and F. J., *Reproducibility of Toolmarks on 20,000 Bullets fired through an M240 Machine Gun Barrel*. AFTE Journal, 2012. **44**(3): p. 248-253.

LaPorte, D., *An Empirical and Validation Study of Breechface Marks on .380 ACP Caliber Cartridge Cases Fired from Ten Consecutively Finished Hi-Point Model C9 Pistols*. AFTE Journal, 2011. **43**(4): p. 303-309.

#### Feature dependence

Further research is necessary to understand the degree of dependence of the different type of information in order to combine them in a logically correct manner.

#### Feature permanence and robustness

Chumbley, L., et al., *Clarity of Microstamped Identifiers as a Function of Primer Hardness and Type of Firearm Action*. AFTE Journal, 2012. **44**(2): p. 145-155.



### 5.2.2.2. Validation of the performance

A large research and development activity has been developed at the NFI to validate the performance of the practitioners. Specific solutions have been implemented in the ACEV protocol in order to manage the context bias and to provide a second and independent opinion. The performance of the practitioners is tested using a system of fake cases and via proficiency tests.

#### Context bias management

Quality document “Context management in firearms comparison casework”

Stoel, R.D., et al., *Minimizing contextual bias in forensic casework*, in *Forensic Science and the Administration of Justice*, SAGE Publishing, H. M. and S. K., Editors: In Press.

#### Second opinion

Quality document “Second opinion procedure”


#### Performance testing

Error rate monitoring using fake cases in which the ground truth is known: quality document “Error protocol in firearms comparisons casework”.

#### Proficiency test

Pauw-Vugts, P., et al., *FAID2009: Proficiency Test and Workshop* AFTE Journal, 2013. 45(2): p. 115-127.

### 5.2.2.3 Assessment of the competence of the practitioner (see further 5.2.2.3)

<b>Summary</b>	Laboratory/customer demands met	
<b>Statement on fitness for intended purpose</b>	 <b>The method fulfills the demand</b>	<input type="checkbox"/> <b>The method does not fulfill the demand</b> <b>The following should be done....</b>