Notice number: 6.001

COMMISSION OF INQUIRY INTO DNA PROJECT 13

Section 5(1)(d) of the Commissions of Inquiry Act 1950

STATEMENT OF GENEROSA LUNDIE

- I, **Generosa Lundie**, care of Queensland Health Forensic and Scientific Service, Forensic Scientist, do solemnly and sincerely declare that:
- 1. On 19 October 2023, I was requested to provide a statement responding to Notice 6.001 "Requirement to Give Information in a Written Statement".

Identification

Question 1(a) - State your full name:

My full name is Generosa Lundie.

Question 1(b) - State your qualifications, skills or experience relevant to forensic science and DNA:

- 3. I graduated from Perpetual Help College of Medicine in the Philippines with a Bachelor of Science Medical Technology in 1991.
- 4. I graduated from Griffith University with a Bachelor of Biomedical Sciences in 2005.
- 5. Annexed and marked Exhibit GL-01 is a copy of my CV.

Question 1(c) - State the period(s) of time you have been or were employed by or otherwise engaged with Queensland Health, Queensland Health Forensic and Scientific Services (QHFSS) and/or Forensic Science Queensland, and in what roles and when.

6. I commenced working as a Laboratory Assistant at Queensland Health Forensic and Scientific Services – DNA Analysis between March and June 2006. This is now known as Forensic Science Queensland. Since then, I have been engaged in various roles with QHFSS and/or Forensic Science Queensland as set out below.

Automation Project Team - June 2006 to late-2008

- 7. Between approximately June 2006 to the end of 2008, I worked in the Automation Project Team as a newly graduated scientist and I was supervised by Thomas Nurthen who was the Project Manager for the team.
- 8. The members of the Automation Project Team were:
 - (a) Thomas Nurthen;
 - (b) Cecilia Iannuzzi;
 - (c) Iman Muharam;



- (d) Vojtech Hlinka; and
- (e) Breanna Gallagher.
- 9. Breanna and I were junior members of the team. The more senior scientists in the team were Thomas, Cecilia, Iman and Vojtech.
- 10. While in the Automation Project Team, my duties as a newly graduated scientist involved:
 - (a) being trained in manual DNA extraction by Cecilia (Breanna was also trained at the same time with me);
 - (b) assisting in the preparation of mock samples to be used in the Project 9 and Project 11 discussed below;
 - (c) undertaking manual DNA extraction; and
 - (d) undertaking other tasks as instructed by Tom who was as the Project Manager of the Automation Project Team and the more senior scientists, being Cecilia, Iman and Vojtech.
- 11. I understood the purpose of the Automation Project Team was to explore opportunities to use automation in the DNA extraction process. I recall there was a plan to incorporate an automated robotic platform so that both new manual and automated DNA extraction methods could be introduced.

Project 9

- 12. In the early part of my time in the Automation Project Team, I recall assisting the senior scientists (Cecilia, Iman and Vojtech) and Breanna with Project 9.
- 13. At the time I joined the Automation Project Team, the wider Forensic Team was using a DNA extraction solution called Chelex. At the time, if I recall correctly, there was a plan to stop using Chelex as a liquid agent to extract DNA because it took longer to process and apparently it could be harmful to scientists.
- A DNA extraction kit is a small box that contains liquid agent solutions necessary for the DNA extraction process.
- 15. Based on my recent review of the Project 9 report, I believe Project 9 involved the evaluation of various commercial DNA extraction kits, which included:
 - (a) trialling six DNA extraction kits (including Chelex, DNA IQ and four others), using mock samples to identify which performed best; and
 - (b) performing manual DNA extraction processes with the different DNA extraction kits.
- 16. DNA IQ was one of the five new extraction kits that was trialled.

17.	Based on my recent review of the Project 9 report, I believe that the related purpose of
	Project 9 was to identify the best extraction kits that could also be used or trialled in an

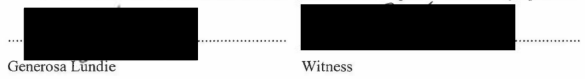
Generosa Lundie	Witness	

automated procedure using an automated robotic platform, being the Multiprobe II platform. The Multiprobe II platform is a robot that automated the DNA extraction process. When I joined the Automation Project Team, there was one or two Multiprobe II platforms being trialled as part of Automation Project Team's work, but I was not involved in that work.

- 18. My involvement in Project 9 occurred a long time ago now, and I only remember bits and pieces such as assisting in the preparation of mock samples and performing laboratory work. I was not trained at that time in DNA analytical processes other than manual extraction. I was not involved in any analysis, evaluation or interpretation of results as I was still new and had limited experience in Forensics.
- 19. I am aware a report was prepared about Project 9.
- 20. I was not involved in interpreting the results or writing the report. I do not know who wrote the Project 9 report. My name is on page 1 of the Project 9 report along with all other members of the Automation Project Team and Vanessa Ientile. I believe all members of the team were named on the report to recognise we all made some wider contribution and to recognise the collective assistance with the work which supported the project.
- 21. **Annexed and marked Exhibit GL-02** is the Project 9 report I found on our digital storage system.

Project 11

- 22. I recall assisting Breanna and the senior scientists with Project 11.
- 23. Based on my recent review of the Project 11 report, I believe this project was about validating that DNA IQ kit (which was identified through Project 9) was the best to use when extracting DNA manually. Again, this took place a long time ago, and I can only recall bits and pieces. I was not involved in any analysis of the results. I assisted in laboratory duties as instructed by the senior scientists.
- 24. I am aware a report was prepared about Project 11. I may have had a brief look at the report before it was finalised, but I do not have a deep knowledge of the data presented in it. As mentioned, I was new to the team with limited experience in Forensics. I do not know who wrote the Project 11 report. I did not have any involvement in drafting the Project 11 report. I believe my name appears on the report to recognise all members of the Project Automation Team made some contribution to the work which supported the project.
- Annexed and marked Exhibit GL-03 is the Project 11 report I found on our digital storage system.
- 26. As a career development task, I asked Tom if I could attend the Australian and New Zealand Forensic and Scientifics Services Symposium in 2008. I asked Tom if I could develop a poster about the outcome of Project 11 for the symposium. Tom agreed, and the poster I developed was approved by Tom and Iman. I displayed the poster at the symposium. After the symposium, the poster was made available to Forensic and Scientific Services staff to inform them about what was presented at the symposium.



Annexed and marked Exhibit GL-04 is a copy of this poster I found on our digital storage system.

Project 13

- 27. I am aware there was a Project 13. I do not recall being involved with this project because between 2006 and 2008 if I recall correctly, I was heavily involved in manual DNA extraction processes. I think Project 13 occurred after Projects 9 and 11 were completed, but some work may have occurred simultaneously.
- 28. I do not recall reviewing or reading any version of the Project 13 report until I was provided with a copy in relation to this Inquiry. I do not know who wrote the Project 13 report. I did not have any involvement in drafting the Project 13 report. I believe my name appears on the report to recognise all members of the Project Automation Team made some contribution to the work which supported the project.
- 29. When I heard in early October 2023 that the Project 13 report was going to be the subject of this Commission of Inquiry, I looked at the report briefly, I did not read it in detail. I saw on pages 8 and 9 that there were highlighted sections and question marks, and this indicated to me that I was looking at a draft report.
- 30. **Annexed and marked Exhibit GL-05** is the Project 13 report I looked at for this purpose, with my markings.

Analytical Team - Late-2008 to August 2023

- 31. I think in late-2008 or early 2009, the Project Automation Team work finished and I was moved to the Analytical Team, where I remained until August 2023.
- 32. I reported to Allan McNeven and my work involved:
 - (a) General laboratory duties;
 - (b) Automated DNA extraction tasks;
 - (c) Manual DNA extraction tasks;
 - (d) Pre-PCR tasks;
 - (e) Capillary Electrophoresis tasks; and
 - (f) Non-laboratory tasks.

Evidence Recovery Team - August 2023 to present

33. In August 2023, I moved to the Evidence Recovery Team. The work in this team involves examination of items, examination of sexual assault kits currently called (FMEKs) and performing presumptive testing on samples under supervision before they are sent to the Analytical section for further testing.

Manual and Automated DNA Extraction Methods

Question 2 - In relation to the report being	the "Project 13. Report on the	Verification of
an Automated/DNA IO Protocol using the N	Jultiprobe ILPLUS HT EX wit	h Gripper
Generosa Lungie	Witness	

Integration Platform", Nurthen, T., Hlinka, V., Muharam, I., Gallagher, B., Lundie, G., Iannuzzi, C., Ientile, V. Automation/LIMS Implementation Project, DNA Analysis FSS (August 2008) (2008 Report)1 and the abstract and introduction therein which state:

1. Abstract

A manual method for extracting DNA from forensic samples using the DNA IQ™ system (Promega Corp., Madison, WI, USA) was validated for routine use in DNA Analysis (FSS). We have verified an automated DNA IQ™ protocol in 96-well format for use on the MultiPROBE® II PLUS HT EX Forensic Workstation platforms (PerkinElmer, Downers Grove, IL, USA). Data indicate that results from the automated procedure are comparable to those from the manual procedure. Contamination checks were performed using samples prepared in checkerboard and zebra-stripe format, and results were as expected. We recommend the use of the MultiPROBE® II PLUS HT EX platforms to perform automated DNA extraction using the DNA IQ™ system.

Introduction

The MultiPROBE® II PLUS HT EX FORENSIC WORKSTATION platforms (PerkinElmer, Downers Grove, IL, USA) are equipped to perform automated DNA extractions, as they include a DPC shaker and individual heat controllers to enable on-board lysis and incubation steps. Currently in DNA Analysis, the MultiPROBE® platforms allow walk-away operation of PCR setup protocols for DNA quantitation and amplification.

The DNA IQ™ protocol has been verified or validated by various laboratories for use on the MultiPROBE® II PLUS platform. The laboratories that perform an automated DNA IQ™ protocol include PathWest (Western Australia). Forensic Science South Australia (South Australia) and Centre of Forensic Sciences in Toronto (Ontario). The MultiPROBE® II PLUS instrument comes pre-loaded with an automated DNA IQ™ protocol. Unlike the other laboratories, however, we did not validate the included protocol, but instead validated a manual DNA IQ™ protocol which was based on the CFS automated protocol (PerkinElmer, 2004), followed by verification of an automated protocol based on the validated manual method.

The verified automated DNA IQ™ protocol is identical to the validated manual protocol used in-house: there are no differences in reagents or volumes. The adopted DNA IQ™ protocol differs slightly, however, from the manufacturer's protocol, as it includes a lysis step using Extraction Buffer (10mM Tris. 1mM EDTA, 100mM NaCl. 20% w/v SDS) in the presence of Proteinase K, before incubating in the DNA IQ™ Lysis Buffer. Furthermore, the lysis incubation conditions were lowered from 70°C to 37°C in order to accommodate extraction of DNA from heat labile materials such as nylon and polyester. In addition, the automated protocol utilises the SlicPrep™ 96 Device (Promega Corp., Madison, WI, USA) for simultaneous processing of samples in a 96-well format.

Manual Method

Question 2(a) describe, with precision, the "manual method" for extracting DNA from forensic samples using the DNA IQTM system referred to in the first line of the Abstract to the 2008 Report (Manual Method), including whether the Manual Method:

- (i) was devised within the QFSS Forensic DNA Analysis laboratory (Laboratory); or
- (ii) was otherwise a modification of an existing manual method (and if so which method),
- 34. The DNA IQ system came in a kit or box containing different chemical liquid solutions used for extracting DNA. I am not able to describe the method with precision without referring to the manual extraction instructions outlined in the relevant written standard operating procedure which we followed at the time. It has been at least 10 years since I performed the procedure for manual extraction using DNA IQ. I can only remember bit and pieces.
- 35. As I recall, the manual method for extracting DNA using DNA IQ involved the following steps:

(a)	Step I – A scientist would receive from the Evidence Recovery Team a tube
	containing a piece of substrate (which might be a piece of cotton or fabric
	collected by police from a crime scene) which might contain a DNA sample. The
	scientist would add a volume of solution 1 (called the extraction buffer) using a

Witness

Generosa Lundie

- pipette. The scientist would then incubate the tube to heat the solution which opens the cell membrane.
- (b) Step 2 The scientist would then remove the tube from incubation and add solution 2 (which is a lysis buffer) and solution 3 (which is a resin containing paramagnetic beads). These solutions were mixed to break down the cells and isolate the DNA. After mixing, the scientist would remove all liquid from the tube using a pipette so the DNA is left attached to the paramagnetic resin beads.
- (c) Step 3 The scientist would then add a volume of solution 4 (which is called the wash buffer) to the tube and mixed it. The scientist then removed all liquid from the tube using a pipette. The DNA which had bound to the resin would remain in the tube. This washing process would be repeated three times so all remaining contaminants or impurities from the DNA were removed and ideally only pure DNA would remain.
- (d) Step 4 The tube containing the DNA was then opened and left to dry inside a extraction cabinet fume hood for a period of time to remove inhibitors such as alcohol.
- (e) Step 5 The scientist would then add solution 5 (which is an elution buffer) to the tube to release DNA from the paramagnetic resin beads. The scientist collected the solution containing the DNA using a pipette and transferred it to another tube. The scientist repeated this process (i.e. solution 5 was added again, and the DNA was collected again). The tube containing the extracted DNA was stored and was then ready for the next stages of processing.
- 36. After the manual extraction process was complete, another member of the Analytical Team would perform the next stages of processing being:
 - (i) DNA quantification.
 - (ii) DNA amplification; and
 - (iii) Capillary electrophoresis,

to produce a DNA profile result which would then be reviewed by the case scientist for DNA comparison.

- 37. I do not know where this manual method was devised.
- 38. I do not know whether it modified an existing manual method.
- 39. I was trained in the manual extraction process by Cecilia between 2006 and 2007 when I commenced in the Automation Project Team. This training involved me:
 - (a) observing Cecilia performing the manual extraction process three times;
 - (b) performing the manual extraction process three times under Celia's supervision;
 - (c) discussing my competence with Cecilia;



- (d) completing a training module, answering both oral and written questions within module;
- (e) being signed-off by Cecilia as competent to perform manual extraction procedure using DNA IQ.

Question 2(b) describe, with precision, the method by which the Manual Method's "routine use" in DNA Analysis (FSS) was validated

- 40. Based on my recent review of the Project 11 report, I understand that Project 11 involved validation of the use of DNA IQ, but my involvement in that project as I recall was limited to performing laboratory duties which was manual extraction processes only.
- 41. I was not involved in the overall validation or the collection and assessment of the outcome of results.

Question 2(c) state whether, and if so how, the Manual Method differed from or otherwise modified the DNA IQTM protocol that was "verified or validated by various laboratories for use on the Multiprobe® II PLUS platform" (as stated in the second line of the second paragraph of the Introduction to the 2008 Report)

42. I do not know the answer to this question. I do not recall being specifically involved in any modified version of using DNA IQ nor do I recall being involved in any verification of the Multiprobe® II PLUS platform.

Question 2(d) state when the Manual Method was so devised

43. I do not know when the manual method was devised. I was only instructed to undertake manual DNA extractions to assist my team members.

Question 2(e) identify those within the Laboratory responsible for devising the Manual Method

44. I do not know who was responsible for devising the manual method.

Question 2(f) state the reason(s) why the Laboratory chose to devise and to implement the Manual Method

45. I do not know.

CFS Automated Protocol

Question 2(g) describe, with precision, the "CFS automated protocol (PerkinElmer, 2004)" (CFS Automated Protocol) referred to in the seventh line of the second paragraph of the Introduction to the 2008 Report

46. I do not know what this is referring to.

Manual DNA IQTM Protocol

Question 2(h) describe, with precision, the " IQ^{TM} Protocol)" referred to in the seventh li	- 1	,
to the 2008 Report, including whether it:		_
		•••••
Generosa Mindie	Witness	, i

- (i) was developed or otherwise supplied by the manufacturer of the MultiPROBE® II PLUS HT EX FORENSIC WORKSTATION platform;
- (ii) was devised within the Laboratory; or
- (iii) was otherwise a modification of an existing Manual DNA IQTM protocol (and if so which method)
- 47. See above at paragraphs [35] to [39].

Question 2(i) describe, with precision, the method by which the Manual DNA IQ™ Protocol was validated

48. I do not know how the manual method using DNA IQ was validated.

Question 2(j) state whether, and if so how, the Manual DNA IQTM Protocol differed from or otherwise modified the DNA IQTM protocol that was "verified or validated by various laboratories for use on the Multiprobe® II PLUS platform" (as stated in the second line of the second paragraph of the Introduction to the 2008 Report)

49. I do not know.

Question 2(k) state when the Manual DNA IQTM Protocol was so devised

50. I do not know when the manual method using DNA IQ was devised.

Question 2(l) identify those within the Laboratory responsible for devising the Manual DNA IQTM Protocol

51. I do not know.

Question 2(m) state the reason(s) why the Laboratory chose to devise Manual DNA IQ™ Protocol

52. I do not know.

Automated DNA IQTM Protocol

Question 2(n) state whether the "automated DNA IQTM protocol" referred to in the first line of the third paragraph of the Introduction to the 2008 Report (Automated DNA IQTM Protocol) is the same as the automated protocol the subject of the 2008 Report. If it is not, then state the reasons why and describe any differences

53. I do not know.

Question 2(0) state whether, and if so how, the Automated DNA IQ™ Protocol differed from or otherwise modified:

(i) the Manual Method;



- (ii) the DNA IQ[™] protocol that was "verified or validated by various laboratories for use on the Multiprobe® II PLUS platform" (as stated in the second line of the second paragraph of the Introduction to the 2008 Report);
- (iii) the CFS Automated Protocol; and
- (iv) the Manual DNA IQTM Protocol
- 54. I do not know.

Question 2(p) state when the Automated DNA IQTM Protocol was so devised

55. I do not know.

Question 2(q) identify those within the Laboratory responsible for devising the Automated DNA IQTM Protocol

56. I do not know.

Question 2(r) state the reason(s) why the Laboratory chose to devise the Automated DNA IQTM Protocol rather than use the manufacturer method

I do not know.

Multiprobe II PLUS HT EX with Gripper Integration Platform (Multiprobe II Device)

Question 3 - State when the Laboratory received the Multiprobe II Device

- 58. I do not know when the laboratory received the Multiprobe II Device.
- 59. The Multiprobe II Device if I recall correctly was in the laboratory when I commenced in June 2006.

Question 4 - For each of the Manual DNA IQ™ Protocol and the Automated DNA IQ™ Protocol, describe, with precision and completeness, what, if anything was done to the device to modify it, including whether any of the manufacturer's factory settings were changed, and if so which ones and how (including but not limited to temperature settings, reagents and volumes)

60. I do not know. Again, my task was very specific and limited to just assisting in laboratory work.

Question 5 - State when the modifications were made

61. I do not know.

Question 6 - Identify those within the Laboratory responsible for the modifications

62. I do not know.

Question / - State the reasons why	the modifications	were made	
Generosa Lundie	Witness		

63. I do not know.

2008 Report

Question 8 - Describe your role in the preparation of the 2008 Report

64. I did not have any role.

Question 9 - Describe the directions you received in relation to the preparation of the 2008 Report, and identify the person or persons from whom you received those directions.

65. I did not receive any directions.

Question 10 - State the substance of the communications (including discussions) that occurred between you, any other authors of the 2008 Report and/or any supervisor or person in a position of management concerning the purpose(s) or intended purpose(s) of the 2008 Report, including by identifying with whom those communications took place and when.

66. I do not recall having any communications about the 2008 Report.

Question 11 - Identify the persons to whom was the 2008 Report was distributed.

67. I do not know.

Question 12 - In relation to the matters stated in the 2008 Report, state:

Question 12(a) - how the conclusion on page 1 of the 2008 Report that "Data indicate that results from the automated procedure are comparable to those from the manual procedure" was reached, including:

- (i) any discussions or communications between any of the named authors of the 2008 Report and any supervisor or person in a position of management in relation to that conclusion or the referenced data; and
- (ii) how that conclusion can be reconciled with the data and figures outlined in part 6.4 of the 2008 Report.
- 68. I do not know. I do not recall being involved in the 2008 Report.

Question 12(b) - how the recommendations summarised on page 18 of the 2008 Report were decided, including by identifying:

- (i) your role in the decision;
- (ii) with whom and when you communicated (including by way of discussion) with any other person in connection with the recommendations and the decision to make them
- 69. I do not know. I do not recall being involved in the 2008 Report.



All the facts and circumstances declared in my statement, are within my own knowledge and belief, except for the facts and circumstances declared from information only, and where applicable, my means of knowledge and sources of information are contained in this statement.

I make this solemn declaration conscientiously believing the same to be true and by virtue of the provisions of the *Oaths Act 1867*.

TAKEN AND DECLARED before me at Brisbane on 24 October 2023



EXHIBITS INDEX

Exhibits Index - Generosa Lundie Statement

Question	Exhibit	Document Title
1(b)	GL-01	GENEROSA Jenny LUNDIE – CV
1(b)	GL-02	Project 9: Report on the Evaluation of Commercial DNA Extraction Chemistries 2007
1(b)	GL-03	Project 11: Report on Validation of a manual method for Extraction DNA using the DNA IQ System, August 2008
1(b)	GL-04	Chemical Validation of a Manual DNA Extraction Method using the DNA IQ System
1(b)	GL-05	Project 13. Report on the Verification of an Automated DNA IQ Protocol using the MultiPROBE II Plus HT ES with Gripper Integration Platform.



GENEROSA Jenny LUNDIE



Career Goals

Currently employed as a Scientist within the Evidence Recovery section of Forensic Biology at Forensic Science Queensland my goal is to continue utilising my scientific skills and experience in a team environment where Forensic DNA examination is the field of expertise.

To continue develop and advance my knowledge, understanding and contribute to the overall goals of the organisation.

Professional Experience

Scientist

August 2003- Current

Evidence Recovery Section

Forensic Biology - Forensic Science Queensland

Responsibilities

- Perform and record in accurate, concise, and contemporaneous manner, the examination of items submitted to Forensic Biology of items, in-tube items, and Forensic Medical Examination kits according to standard operating procedure.
- · Entry and review of results examination into laboratory information system.
- Participate in internal and external proficiency testing as required.
- Maintain QIS event, participate in training and complete training modules.
- · Provide training to scientists, technician or other staff as directed by the senior scientist.
- Maintenance of scientific knowledge and awareness of new/emerging technologies.

Scientist

2008-2003

Analytical Section

Forensic DNA Analysis- Forensic and Scientific Services Health Support Queensland

Responsibilities

- Perform DNA extraction of casework and reference samples both automated and manual procedures using DNA IQ System and casework Pro kit for Maxwell 16.
- Perform DNA Quantification of extracted DNA using the Quant Studio Real-Time PCR System.
- Amplification of DNA extracts using PowerPlex21 on the Proflex thermal cyclers.
- Perform plate preparation of PCR products for capillary electrophoresis using 3500xl Genetic Analyzer.
- Interpretation of DNA profiles and CE Quality checking using Gene Mapper -- IDX software for PowerPlex21.
- Import/upload results of reference and casework through Forensic Register.
- To ensure consistent production of quality results and assist in the maintenance of quality systems, monitor and update quality systems for example (raising of OQI's).
- Review, manage and validate results of samples and order rework strategy to obtain the best DNA profiles.
- · Assist in the evaluation, verification and validation of new reagents, instruments, and equipment.
- Assist in the training of new staff member, when necessary, in analytical section.
- Perform maintenance and calibration of laboratory equipment and instrument such QIASymphony and Hamilton STARlet liquid handlers.
- Participate in a moot court training session
- · To participate in staff and team meetings and take part in Performance Appraisal Development.
- Perform post-extraction processing of DNA extracts including Microcon, Nucleospin clean-ups, dilutions, and transfers and pooling.

1

- Make standards for quantification of DNA extracts and testing of Quantifiler kits as well.
- Carry out weekly stock take and monthly environmental cleaning as well as general maintenance and cleaning of laboratory.
- Maintaining and updating relevant SOP's and all training requirements are up to date.
- Assist other team member in analytical area if needed, provided all the work required for the day has been completed.
- Ensure there's sufficient batches prepared both manual and automated procedures. Daily recording of Key Performance Index (KPI) for rostered area.
- Keep up to date of laboratory activities and changes by accessing meeting minutes, reading e-mail, Forensic Register Notification, AUSLAB system messages and newsletters.

Scientist

2006-2008

Automation

Forensic DNA Analysis- Forensic and Scientific Services Health Support Queensland

Responsibilities

- Assist in the evaluation and validation of new DNA extraction kits for manual DNA extraction.
- · Assist in the preparation of mock samples to be use in the validation of DNA extraction kits
- Perform operational procedures on the MP II Plus robotic platform including maintenance, gravimetric analysis, deck calibrations and setting-up for extraction processes.
- Use AUSLAB to register and batch functionality.
- Train and supervise staff in the operation of MPII Plus robotic platform using DNA IQ ™ system.
- Participate in team meetings.

Laboratory Assistant

March-June 2006

Operational Team

Forensic DNA Analysis- Forensic and Scientific Services Health Support Queensland

- Responsibilities
- Perform FTA punching & washing to purify and amplify reference DNA samples.
- Perform blood clothing from reference and evidence sample received in the laboratory.
- Utilised AUSLAB for registration, storing, transfer of samples and batch functionality.
- Access QIS (Quality Information System) for updated protocols, QIS notifications and OQI (Opportunity for Quality Improvements).
- Assist in the environmental sampling and cleaning.
- · To participate in staff and team meetings and take part in Performance Appraisal Development.

EDUCATIONAL QUALIFICATION

Bachelor of Biomedical Science

January 2003 - December 2005

Griffith University

Key Subjects

- Cell and Molecular Biology
- Biochemistry
- Laboratory Biotechniques
- Chemistry
- Biological Chemistry

Bachelor of Science in Medical Technology

June 1985 - March 1991

Perpetual Help College of Medicine (Philippines)

Key subjects

- Anatomy and Physiology
- Serology
- Hematology
- Parasitology
- Microbiology and Virology
- Immunology

Trainings

- Completion of the Perkin Elmer Users training for MPII Robotic Platform Workstation including (hardware overview, basic WinPREP Software, applications, specific training, customer preventive maintenance and good liquid handling.)
- Completed the training module in Interpretation and Acceptance of Results using AmpFISTR Profiler and COfiler System™.
- Completed the training of Automated DNA IQ™ Method of Extracting DNA from Reference and Casework Samples.
- Completed the training in "Court Training" and Attendance at both Supreme and Magistrate Court.
- Processing FTA Paper™ using Profiler Plus.
- · AUSLAB Functionality (Laboratory Information and Management System).
- · Real Time PCR Quantification of Human DNA using Quantifiler Human DNA Quantification Kit.
- PCR Amplification using Profiler Plus and COfiler System™.
- Competent to Train in the following modules (Operation, Maintenance and Gravimetric Analysis of the MPII Plus and Automated DNA IQ Method of Extracting DNA from Reference and Casework Samples using DNA IQ Kit.

SKILLS and ATTRIBUTES

Scientific knowledge and skills

Gained through the completion of two medical science degrees, employment as a Scientist in Forensic DNA Analysis and as Medical Technician at Perpetual Hospital.)

Problem solving/trouble shooting

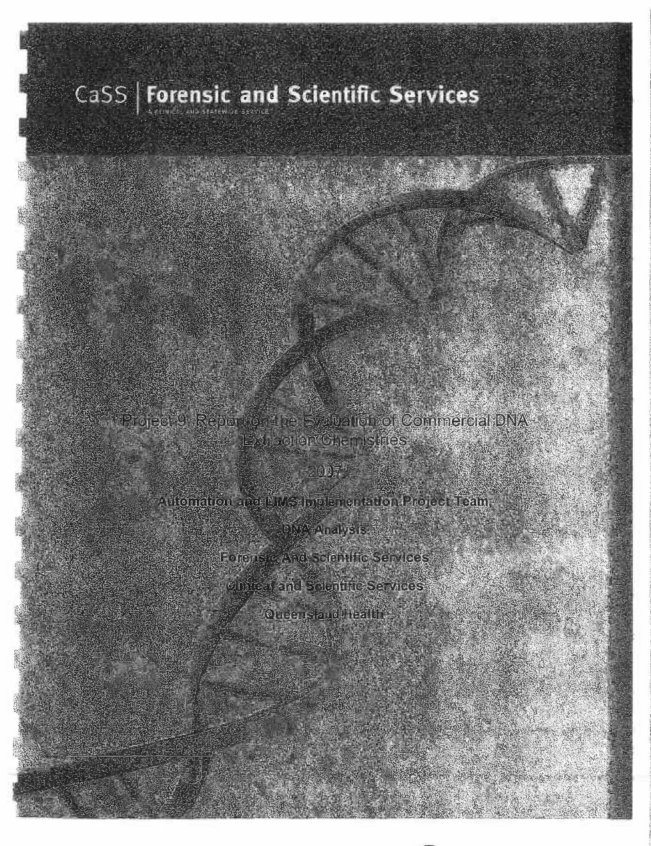
Developed through my involvement in various projects in DNA Analysis for which I participated/assisted in the validation to develop a new method of improving existing working practices.

Effective oral and written communications

Developed through regular interaction, meetings, and seminars attendance both internal and external. Assist in writing reports and updating Standard Operating Procedures.

Excellent teamwork skills

Through participation in team meetings, share information, contribute to all activities, and work co-operatively with team members to improve and achieve team performance and objectives.



safe i sustainable i appropriate



Project 9. Report on the Evaluation of Commercial DNA **Extraction Chemistries**

Breanna Gallagher*, Vojtech Hilnka*, Cecifia lannuzzi*, Generosa Lundle*, Iman Muharam*, Thomas Nurthen

^ These authors contributed equally.

Automation/LiMS implementation Project, DNA Analysis FSS (June 2007)

1. Abstract

DNA Analysis FSS performed an evaluation of various commercial DNA extraction chemistries in order to compare their overall performance (quality, yield user friendliness and the ability to automate) against the current in-house Chelex® profocol. Out of five commercial kits evaluated, the DNA IQ™ system from Promega Corporation (Madison, Wi, USA) was found to be the best out-of-the-box method for DNA extraction of blood and cell samples and will be validated for routine in-house use. This document presents data from the evaluation and provides a discussion of the results observed. For the manual DNA IQ™ validation report, see Project 11. Verification of an automated DNA IQ method is reported in Project 13.

2. Introduction

There have been many DNA extraction methods published since DNA was first isolated in 1953 (Butler, 2005). As technology developed and the demand for DNA testing there as the methods for extracting and purifying DNA have improved. The Chelex extraction procedure (Walsh et al., 1991) became a quick and easy alternative to the more technically demanding phenol/chloroform protocol and was more compatible for extracting samples from forensic exhibits, although the resulting DNA extract is still cruce and unpurified because inhibitors are not removed from the solution. As the demand for extracting trace DNA samples has increased within the last 10 years to allow interrogation. of low copy number forensic samples, coupled with the increase in the need to analyse difficult samples such as touched objects and degraded bone material, new DNA extraction technologies that are designed specifically for forensic samples have increased in

The new DNA extraction chemistries on the market aim to overcome problems encountered in forensic DNA samples as they are designed to:

- Improve removal of inhibitors present in the sample that can affect DNA extraction (e.g. hemoglobin, textile dyes) or prevent successful PCR amplification (e.g. hematin, melanin, polysaccharides, bile salts, humic compounds);
- Maximise recovery of DNA in trace (low copy number) samples by using special buffers that promote cell lysis and integrating a DNA capture system that allows efficient binding and elution of sample DNA, therefore increasing total yields;
- Increase the overall quality and purity of recovered DNA by using special elution of storage buffers, therefore enhancing DNA stability for long-term storage, ensuring reliability and consistency in the sample DNA for reworks and future use.

DNA Analysis FSS obtained various commercial forensic DNA extraction kits (Table 1) in order to evaluate their performance against the in-house Chelex® protocol (see QIS 17171 for detailed information and literature on the Chelex® system).



Table 1. Extraction kits that were evaluated by Forensic Biology FSS.

DNA extraction kit and manufacturer

DNA IQ™ (Promega Corp., Madison, WI, USA) DNA IQ ™ (Promega Corp., Madison, WI, USA)
QIAamp® DNA Micro (Qiagen GmbH, Hilden, Germany)
ChargeSwitch® (Invitrogen, Carlsbad, CA, USA)
forensicGEM™ (ZyGEM, Hamilton, NZ)
NucleoSpin® 8 Trace (Macherey-Nagel, Düren, Germany)

Technology type

Novel paramagnetic beads Silica-based membrane Magnetic beads Thermophilic proteinase incubation Silica-based membrane

Magnetic bead technology is based on the use of magnetic resin that has the capability bind DNA when subjected to a particular environmental pH or lonic strength. Therefore, by using buffers with different pH values or different ionic components, the binding and elution of DNA can be controlled. Furthermore, whilst the DNA is bound to the resin DNA complex can be washed using an alcohol-containing buffer in order to remove inhibitors. and residual proteins. A magnetic force is applied during the washing procedure to immobilise the resin-DNA complex and ensure no DNA is lost during washing. Membrane technology is based on a similar principle, except the DNA is immobilised in a thin silicabased membrane within the column.

forensicGEM™, the recently-released one-tube proteinase incubation system uses a thermostable enzyme to digest nucleases in order to yield a crude DNA extract. The enzyme digest method does not incorporate any washing steps, however, and therefore inhibitors are not removed from solution.

3. Aim

To evaluate several commercial DNA extraction kits (as per Table 1) that were specifically designed for forensic DNA samples, using the manufacturer's recommended manual protocols, and compare against the current in-house Chelex® protocol, in order to select a sultable kit for manual validation and automated verification.

4. Equipment and Materials

- Chelex®-100, P/N 143-2832 (Biorad, Hercules, CA, USA) DNA IQ™ System, P/N DC6701 (Promega Corp., Madison, WI, USA)
- QIAamp® DNA Micro Kit, P/N 56304 (Qiagen GmbH, Hilden, Germany)
- ChargeSwitch® Forensic DNA Purification Kit, P/N CS11200 (Invitrogen, Carlsbad CA, USA)
- forensicGEM™ (ZyGEM, Hamilton, NZ)
- NucleoSpin® 8 Trace, P/N 740 722.1 (Macherey-Nagel, Düren, Germany)

For preparation of buffers and reagents specific for each kit, see the Methods section that is relevant for that kit.

5. Methods

5.1 Mock sample creation

Refer to document "Mock sample creation for cell and blood samples" (Gallagher et al., 2007) for the detailed protocol.



5.2 DNA extraction kit protocols

The following section provides the principle and protocol for each DNA extraction kit as recommended by the manufacturer. The Chelex® method was as per QIS 17171.

5.2.1. Chelex®-100 (BioRad)

Principle

Chelex® is a chelating resin composed of styrene divinylbenzene copolymers which have a high affinity for polyvalent metal ions. The copolymers contain paired iminodiacetate ions acting as chelating groups which chelate metal ions, including some that degrade DNA while boiling the sample to obtain elitied DNA. Chelex is the current Forensic Biology FSS standard in-house extraction protoco

- Equipment and Materials
 o 20% Chelex® solution (w/v)
 - Waterbath
 - Magnetic stirrer plate
 - 1.5mL sterile tubes
 - Spin baskets
 - Autoclaved nanopure water
 - Vortex
 - Centrifuge
 - Twirling sticks

 - Proteinase K (10mg/mL) FTA® Classic Card, P/N WB120205 (Whatman Plc)

Preparation of reagents

20% Chelex®-100

On balance, to a beaker containing a magnetic stirrer bargadd 2 grams of Chelex®-100 resin. To this, add 10mL of autoclaved hanopure water to make a 20% w/v solution and cover with parafilm. To ensure that the Chelex® is evenly dispersed, place beaker onto a magnetic stirrer plate before pipetting.

Methods (see QIS 17171R9)

- Label sterile 1.5mL screw-capped tubes which contain sample as well as new elution tubes including extraction controls.
- Pipette 1mL of autoclaved nanopure water into each tube, vortex gently.
- incubate at room temperature for 30 minutes.

The following steps are determined by sample type:

- For buccal FTA® punches, place tubes on multitube vortex for 5min at 12,000rpm.
- For cell and/or fabric samples, twirl the substrate with a sterile twirling stick

Note: Vortex FTA® punches samples then go to "For all sample types."

- Transfer swab/fabric into spin baskets.
- Spin tubes with spin basket for 30s at maximum speed (~15,800g or the applicable centrifuge's maximum setting). Discard spin basket with swab.



8. Vortex supernatant, then pour back into original extract tube.

For all sample types

- Vortex, then spin in centrifuge for 3min at maximum speed (~15,800g or the applicable centrifuge's maximum setting).
- Carefully remove all but 50µL of supernatant. Leave substrate in tube with pellet.
- 11. Add 150µL of 20% Chelex® to each tube and vortex.

Note: When pipetting Chelex, the resin beads must be distributed evenly in the solution. Use magnetic stirrer in beaker of Chelex and wide bore pipette tips.

- 12. Add 4μL of Proteinase K (10mg/mL) to cells and mix gently by vortexing
- 13. Incubate in 56°C water bath for 30min for blood and cell samples.
- 14. Vortex until mixed, then incubate in boiling water bath for 8min.
- Vortex until mixed, then centrifuge for 3min at maximum speed (\$15,800g or the applicable centrifuge's maximum setting).
- Transfer supernatant to new labelled 1.5mL screw-capped tube leaving Chelex® beads behind.
- 17. Samples are stored at -20°C.

5.2.2. DNA IQ™ System (Promega Corp.)

Principle

The Promega DNA IQ™ system for small casework samples incorporates two distinct steps. The first step provides an easy, rapid, efficient and almost universal cell lysis method to extract biological materials off stains on solid supports. The second step utilised a specific paramagnetic resin that purifies DNA without extensive washing to remove the lysis reagent. The DNA IQ™ system is designed to purify DNA samples approximately 100ng or less, and is more efficient with samples containing less than 10ng of DNA.

Equipment and Materials

- DNA IQ™ System (100 samples, Cat.# DC6701) containing:
 - o 0.9mL Resin
 - o 40mL Lysis Buffer
 - o 30mL 2X Wash Buffer
 - o 15mL Elution Buffer
- MagneSphere® Magnetic Separation Stand, 12-position (Cat.# Z5342)
- o DNA IQ™ Spin Baskets (Cat.# V1221)
- o Microtube 1.5mL (Cat.# V1231)
- o 95-100% ethanol
- o Isopropyl alcohol
- o 1M DTT
- o 65°C heat block
- o 70°C heat block
- Vortex mixer

Preparation of Buffers

- Preparing 1X Wash Buffer
 - For DC6701 (100 samples), add 15mL of 95-100% ethanol and 15mL of isopropyl alcohol to 2X Wash Buffer.
 - ii. Replace cap and thoroughly mix by inversion.



age 4 of 34

- iii. Mark label to record addition of alcohols.iv. Label bottle as "1X Wash Buffer".
- Store bottle at room temperature with lid closed tightly to prevent evaporation.

Preparing Lysis Buffer

i. Determine the total amount of Lysis Buffer to be used (Table and add 1µL of 1M DTT for every 100µL of Lysis Solution,

Table 2. Total amount of Lysis Buffer required for different sample material type

Material	Lysis Buffer	Lysis Buffer	Total Buffer
Liquid blood	100µL	100µL 388	€ ₹200µL
Cotton swab	250µL	100µL	350μ
1/4 th CEP swab	250µL	100 µL	350µL
15-50mm2 S&S 903 paper	150µL	100µL	250µL
3-30mm2 FTA® paper	150µL	100µL.迎常	250µL
Cloth up to 25mm ²	150ul-	10011	250 190

For use in Step 2; For use in Step 9.

- ii. Mix by inversion.
- iii. Mark and date label to record addition of DT
- iv. Seal tube and store solution at room temperature for up to one month if required.

DNA isolation from stains on solid material (non-liquid samples)

- Place sample in a 1.5mL Microtube. The recommended amount of resin can capture a maximum of ~100ng DNA, therefore consider this when determining amount of sample to add.
- Add 250µL of prepared Lysis Buffer (Table 2). Close lid and place on a 70°C heat block for 30min.
- Remove tube from heat block and transfer the Lysis Buffer and sample to a DNA IQ™ Spin Basket.
- Centrifuge at room temperature for 2min at maximum speed. Remove spin basket.
- Vortex the stock Resin for 10s until it is thoroughly mixed. Add 7µL Resir to the sample. Keep the Resin resuspended while dispensing to obtain uniform results.
- Vortex sample / Lysis Buffer / Resin mix for 3s. Incubate at room temperature for 5min.
- Vortex for 2s and place tube in the MagneSphere® Magnetic Separation Stand, Separation will occur instantly.
- Carefully remove and discard all of the solution without disturbing the Resin on the side of the tube.
- Add 100µL of prepared Lysis Buffer. Remove the tube from the MagneSphere Magnetic Separation Stand and vortex for 2 seconds.
- 10. Return tube to the MagneSphere® Magnetic Separation Stand and discard all Lysis Buffer, without disturbing the resin on the side of the tube.
- 11. Add 100µL prepared 1X Wash Buffer. Remove tube from the MagneSphere Magnetic Separation Stand and vortex for 2s.
 12. Return tube to the MagneSphere Magnetic Separation Stand and discard
- all Wash Buffer, without disturbing the resin on the side of the tube.
- 13. Repeat steps 11 and 12 once for a total of 2 washes. Make sure that all of the solution has been removed after the last wash.



CaSS Forensic and Scientific Services

- 14. With lid open, air-dry the Resin in the MagneSphere® Magnetic Separation Stand for 5min to 15min.
- 15. Add 25-100µL Elution Buffer, depending on how much biological material was used. A lower elution volume ensures a higher final concentration of DNA.
- 16. Close the lid, vortex the tube for 2s and incubate at 65°C for 5mig
- 17. Remove the tube from the heat block and vortex for 2s. Immediately blac on the MagneSphere® Magnetic Separation Stand.
- 18. Transfer the solution to a fresh tube.
- 19. Store the DNA extract at 4°C for short-term storage or at 20 or 170°C long term storage.

5.2.3. QlAamp® DNA Micro (Qiagen)

Principle

The QIAamp® DNA Micro kit combines selective binding properties of a silica-based membrane with flexible elution volumes that is suitable for a wide range of sample materials such as small volumes of blood, blood eards, small tissue samples and forensic samples. The basic procedure consists of 4 steps:

- Lysis: the sample is lysed;
- Bind: the DNA in the lysate binds to the membrane of the QIAamp MinElute column;
- Wash: the membrane is washed;
- Elute: DNA is eluted from the membrane.

Equipment and Materials

- QIAamp® DNA Micro kit containing:

 O QIAamp® MinElute Columns;

 - collection tubes (2mL);
 - Buffer ATL;
 - Buffer AL;
 - Buffer AW1 (concentrate);
 - Buffer AW2 (concentrate);
 - Buffer AE;
 - carrier RNA (red cap);
 - Proteinase K.
- Ethanol (96-100%)
- 1.5mL or 2mL microcentrifuge tubes (for lysis steps)
- 1.5mL microcentrifuge tubes (for elution steps)
- Thermomixer
- Microcentrifuge with rotor for 2mL tubes
- Scissors
- Blood collection cards or FTA® card
- Sterile cotton swabs

Important points before starting

- Perform all centrifugation steps at room temperature (15-
- Check whether carrier RNA is required; for purification of DNA from very small amounts of sample, such as low volumes of blood (<10µL) or forensic samples, it is recommended to add



carrier RNA to Buffer AL. For samples containing larger amounts of DNA, addition of carrier RNA is optional.

Steps to perform before starting

- Equilibrate Buffer AE or distilled water for elution to room; temperature (15-25°C).
- Set a thermomixer or heated orbital incubator to 56°C for use in step 2, and a second thermomixer or heated orbital incubator to 70°C for use in step 5. If thermomixer or heated orbital incubators are not available, heating blocks or water baths can be used instead.
- If Buffer AL or Buffer ATL contains precipitates dissolve by heating to 70°C with gentle agitation.
- If processing semen stains, hair, or nathelippings, prepare an aqueous 1M DTT (dithiothreitol) stock solution. Store aliquots at -20°C. Thaw immediately before use:
- Ensure that Buffers AW1 and AW2 have been prepared according to the instructions.

Preparation of Buffers

- Preparing Buffer ATL
 Before starting the procedure, check whether precipitate has formed in
 Buffer ATL. If necessary, dissolve by heating to 70°C with gentle a
 gitation
- Preparing Buffer AL
 Before starting the procedure, check whether precipitate has formed in
 Buffer AL. If necessary, dissolve by heating to 70°C with gentle
 agitation.
- Preparing Buffer AW1
 Add 25mL ethanol (96-100%) to the bottle containing 19mL Buffer AW1 concentrate. Tick the check box on the bottle label to indicate that ethanol has been added. Reconstituted Buffer AW1 can be stored at room temperature (15-25°C) for up to 1 year. Note: before starting the procedure, mix the reconstituted Buffer AW1 by shaking.
- Preparing Buffer AW2
 Add 30mL ethanol (96-100%) to the bottle containing 13mL Buffer AW2 concentrate. Reconstituted Buffer AW2 can be stored at room temperature (15-25%) for up to 1 year. Note: before starting the procedure, mix the reconstituted Buffer AW2 by shaking.

Method

- Lysing material stained with blood or saliva: cut out up to 0.5cm² of stained material and then cut into smaller pieces. Transfer the pieces to a 2mL microcentrifuge tube. Add 300μL buffer ATL, and 20μL Proteinase K. Close the lid and mix by pulse-vortexing for 10s. Continue this procedure from step 2.
- Place the tube in a thermomixer or heated orbital incubator, and incubate at 56°C with shaking at 900rpm for at least 1hr. In general, hair is lysed in 1hr. If necessary, increase the incubation time to ensure complete lysis.
- 3. Briefly centrifuge the tube to remove droplets from the inside of the lid.
- Add 300µL Buffer AL, close the lid, and mix by pulse vortexing for 10s. To
 ensure efficient lysis, it is essential that the sample and buffer AL are



Page 7 of 34

thoroughly mixed to yield a homogeneous solution. A white precipitate may form when Buffer AL is added to buffer ATL. The precipitate does not interfere with the QIAamp® procedure and will dissolve during incubation in step 5. Note: if carrier RNA is required, add 1µg dissolved carrier RNA to 300µL buffer AL.

 Place the tube in the thermomixer or heated orbital incubator, and incubates, at 70°C with shaking at 900rpm for 10min. If using a heating block or water bath, vortex the tube for 10s every 3min to improve lysis

 Centrifuge the tube at full speed on a bench top centrifuge (20,000g 14,000rpm) for 1min.

7. Carefully transfer the supernatant from step 6 to the Qiaamp® Mintipute column without wetting the rim. Close the lid, and centarge at 6,000g (8,000rpm) for 1min. Place the QIAamp® Mintipute column in a clean 2mL collection tube, and discard the collection tube containing the flow through

 If lysate has not completely passed through the membrane after centrifugation, centrifuge again at a higher speed until QIAamp[®] MinElute

column is empty.

Carefully open the QIAamp[®] MinElute column and add 500µL Buffer WA1 without wetting the rim. Close the Ild and centrifuge 6,000g (8,000rpm) for 1min. Place the QIAamp[®] MinElute column in a clean 2ft collection tube and discard the collection tube containing the flow-through.
 Carefully open the QIAamp[®] MinElute column and add 500µL Buffer AW2

10. Carefully open the QIAamp® MinElute column and add 500µL Buffer AW2 without wetting the rim. Close the lid and centrifuge at 6,000g (8,000rpm) for 1min. Place the QIAamp® MinElute column in a clean 2mL collection tube, and discard the collection tube containing the flow-through. Contact between the QIAamp® MinElute column and the flow-through should be avoided. Some centrifuge rotors may vibrate upon deceleration, resulting in the flow through, which contains ethanol, coming into contact with the QIAamp® MinElute column. Take care when removing the QIAamp® MinElute column and collection tube from the rotor, so that flow-through does not some into contact with the QIAamp® MinElute column.

does not come into contact with the QIAamp® MinElute column.

11. Centrifuge at full speed (20,000g; 14,000rpm) for 3min to dry the membrane completely. This step is necessary, since ethanol carryover into

the eluate may interfere with some downstream applications.

12. Place the QIAamp® MinElute column in a clean 1.5mL microcentrifuge tube and discard the collection tube containing the flow through. Carefully open the lid of the QIAamp® MinElute column and apply 45µL Buffer AE (equilibrated to room temperature) to the centre of the membrane to ensure complete elution of bound DNA. QIAamp® MinElute columns provide flexibility in the choice of elution volume.

13. Close the lid and incubate at room temperature (15-25°C) for 1min. Centrifuge at full speed (20,000g; 14,000rpm) for 1min. Incubating the QIAamp® MinElute columns loaded with Buffer AE or water for 5min at room temperature before centrifugation generally increases DNA yield.

5.2.4. ChargeSwitch® (Invitrogen)

Principle

ChargeSwitch® uses a novel magnetic bead-based technology known as ChargeSwitch Technology® (CST®). CST® provides a switchable surface charge, which is switched on and off by changing the pH. With a low pH buffer, the negatively charged DNA backbone binds to the positively charged beads and with a high pH buffer, DNA is eluted by neutralising the charge on the beads.



Page 8 of 34

ChargeSwitch® uses a universal lysis step for all forensic sample types and has been designed to elute DNA from small sample volumes.

ChargeSwitch® uses a basic 4 step principle:

- Lyse sample:
- Negatively charged DNA binds to positively charged beads in a buffer with a pH ≤ 6 so charge is switched on;
- At a pH of 7, charge is still on while beads and bound DNA is washed removing any contaminants;
- In a buffer with a pH of 8.5, charge is switched off and DNA is cluted from the beads.

- Equipment and Materials

 OhargeSwitch® Forensic DNA Purification kit (signed at room)

 - temperature) includes (for 100 preps):

 o ChargeSwitch® Lysis Buffer (L13) 100mL
 o ChargeSwitch® Magnetic Beads (storage buffer::10mM MES, pH
 - 5.0, 10mM NaCl, 0.1% Tween 20) 2 x 1ms Proteinase K (20mg/ml in 50mM Tris-HCl, 11 8.5, 5mM CaCl²

 - 50% glycerol stored at 4°C) 1mL
 ChargeSwitch® Purification Buffer (N5) 20mL
 ChargeSwitch® Wash Buffer (W12) 100mL
 ChargeSwitch® Elution Buffer (E5; 10mM Tris-Hel, pH 8
 - MagnaRack™, P/N CS15000 (Invitrogen)
 - Sterile, 1.5mL microcentrifuge tubes
 - Vortex mixer
 - Waterbath set at 55°C

- Set water bath at 55°C and prepare Lysis master mix in appropriate sized tube using the following formula: n x (1mL ChargeSwitch® Lysis buffer + 10μL Proteinase K) where n is the number of samples.
- To tube add 1mL of ChargeSwitch® Lysis Buffer (L13) and immerse forensic sample in mix.
- Vortex/invert samples for 10-15s to mix then incubate in 55°C water bath for 1hr. Incubation can be shortened to 30min if sample is vortexed or inverted during this step.
- Remove sample or transfer lysate to clean tube using 1mL pipette tips and pipette.
- Vortex ChargeSwitch® Magnetic Beads to resuspend evenly in storage
- Add 200µL of ChargeSwitch® Purification Buffer (N5) to lysate and mix gently by pipetting up and down.
- Add 20µL of ChargeSwitch® Magnetic Beads to sample. Pipette-mix to ensure that no bubbles form.
- 8. Incubate for 1-5min at room temperature to allow the DNA to bind and then place sample tube in MagnaRack™ until a tight pellet has formed. Once this has occurred, aspirate supernatant from tube whilst still in rack and discard, ensuring that the pellet is not disturbed.
- When supernatant has been completely discarded, remove tube from rack and add 500µL ChargeSwitch® Wash Buffer (W12). Mix gently by pipetting up and down to resuspend the pellet.



- 10. Allow beads to form a tight pellet by placing tube in MagnaRack™ and remove supernatant completely, without removing from rack or disturbing the pellet and discard.
- 11. Repeat steps 9 and 10 again.
- 12. Remove tube from rack, ensuring that supernatant has been completely, removed and add 150µL ChargeSwitch® Elution Buffer (E5). Mix.by pipetting up and down 10 times.
- 13. At room temperature, incubate for 1-5min then resuspend peller and min like In step 12.
- 14. Place tube in MagnaRack™ for 1min or until a tight pellerforms. Without removing tube from rack, aspirate DNA supernatant and place in a sean, sterile 1.5mL microcentrifuge tube, ensuring that the pellet is not disturbe If elution is discoloured repeat steps 12 to 14 again
- 15. Discard beads once extraction process is finished and either quant immediately or store at -20°C.

5.2.5. forensicGEM™ (ZyGEM)

Principle

forensicGEM™ is a novel thermophilic proteinase developed as a lapid, cheap and effective DNA extraction solution for forensic laboratories that was recently released. It is a simple closed tube forensic DNA extraction method using a thermostable proteinase.

Protocols are available for blood and cell samples.

Equipment and Materials o forensicGEMTM buffer

- forensicGEMTM
- Heat block or water bath set at 75°C and 95°C
- 20µL sterile Aerosol Resistant Tips
- 0.5-10µL pipettor
- 300µL sterile Aerosol Resistant Tips
- 20-200µL pipettor
- 1mL sterile Aerosol Resistant Tips
- 50µL-1mL pipettor

Method

DNA extraction from buccal swabs using forensicGEMTM

- Add buccal swab to tube.
- Note: 1/4 head of swab specified but can utilise up to whole swab. Add 200µL of forensicGEMTM buffer. Note: if more than 1/4 head of buccal swab is used need to add more forensicGEM buffer. Moss et al. (2003) added 200µL more of the
- forensicGEM buffer for trace samples.

 Add 2µL of forensicGEMTM buffer and forensicGEMTM.

 Note: forensicGEMTM buffer and forensicGEMTM can be added as a mastermix.
- Incubate at 75°C for 15min.
- Incubate at 95°C for 5min.
- Remove supernatant to a new tube for storage.

DNA extraction from FTA® containing blood or salive using forensicGEM™

UV irradiate plasticware for 5min.



- Add FTA® punches to each well of a 96-well plate. Note: Larger punches can be added but not scalable SOP. PCR tubes can also be used for processing.
- Add 100µL H₂O and leave at room temperature for 15min.
- Decant water (remove by pipetting).
 Add 100µL forensicGEMTM buffer and 2µL of forensicGEMTM Note: The method is not listed as scalable.
- Incubate at 75°C for 15min.
- Incubate at 95°C for 5min.
- Remove supernatant to a new tube for storage.

5,2.6. NucleoSpin® 8 Trace (Macherey-Nagel)

Principle

With the NucleoSpin® 8 Trace method, genomic DNA is prepared from forensic samples. Lysls is achieved by incubation of samples in a solution containing. chaotropic ions in the presence of proteinase K at room temperature. Appropriate conditions for binding of DNA to the silica membrane in the NucleoSpin Prace Binding Strips are created by addition of isopropanol to the lysate. The binding process is reversible and specific to nucleic acids. Inhibitors are removed by two washing steps with ethanolic buffer. Pure genomic DNA is finally eluted under low ionic strength conditions in a slightly alkaline elution buffer.

- Equipment and Materials

 o NucleoSpin® 8 Trace kit, containing:
 o Buffer FLB

 - Buffer B5 (concentrate)
 - Proteinase K (lyophilised)
 - Proteinase Buffer
 - Buffer BE
 - NucleoSpin® Trace Binding Strips
 - MN Wash Plate
 - MN Square-well Blocks
 - MN Tube Strips
 - Cap Strips
 - Self-adhering PE Foil
 - NucleoSpin® 8 Trace Starter Set A containing Column Holders A and Dummy Strips
 - PVM vacuum manifold (from MultiPROBE® II PLUS HT EX platform)

Preparation of Buffers

- Proteinase K Add 3mL Proteinase Buffer per vial to dissolve the lyophylised proteinase K and store at -20°C.
- Add 160mL ethanol to 40mL Buffer B5.
- Store all other components of the kit at room temperature. Storage at lower temperatures may cause precipitation of salts. If a salt precipitate is observed, incubate the bottle at 30-40°C for a few minutes and mix well until all precipitation is redissolved.



Method

- Premix 25µL Proteinase K and at least 125µL buffer FLB and add to sample. Incubate the sample at room temperature for 3 hours.
- Insert spacers "MTP/Multi 96 plate" into the vacuum manifold. Place the waste container inside the vacuum manifold and insert a MN Wash Plate into the notches of the spacers. Close the manifold with the lid.
- Place a NucleoSpin® Trace Binding Strips inserted in Column Holder A Into the rubber seal of the vacuum manifold's lid and apply the samples to the wells of the plate.
- Add 1 volume isopropanol to 2 volumes of lysate, mix three times and transfer to NucleoSpin® Trace Binding Strips.
- Bind genomic DNA by applying vacuum until all lysates have passed through the columns (-200mbar 2min; -600mbar 10s). Ventilate the vacuum manifold.
- Wash silica membrane by adding 900µL Buffer B5 to each well of the NucleoSpin® Trace Binding Strips. Apply vacuum (200mbar 1min) until all buffer has passed through the columns. Ventilate the vacuum manifold.
- 7. Repeat the wash procedure once.
- After the final washing step, close the valve, ventilate the vacuum manifold and remove the wash plate and waste container from the vacuum manifold.
- 9. Remove any residual washing buffer from the NucleoSpin Trace Binding Strips. If necessary, tap the outlets of the NucleoSpin Trace Binding Strips onto a clean paper sheet (supplied with the MN Wash Plate) or soft tissue until no drops come out. Insert the column holder with NucleoSpin Trace Binding Strips into the lid and close the manifold. Apply maximum vacuum (-600mbar) for at least 10min to dry the membrane completely This step is necessary to eliminate traces of ethanol. Close the valve and ventilate the vacuum manifold.
- 10. For elution, insert spacers "Microtube Rack" into manifold and rest rack with MN Tube Strips on spacers. Insert Column Holder A with NucleoSpin Trace Binding Strips into manifold lid. Pipette 100µL Buffer BE directly to the bottom of each well and incubate for 5min at room temperature. Apply vacuum (-400mbar 2min).

5.3 DNA quantitation

All DNA extracts were quantified using the Quantifiler™ Human DNA Quantitation kit (Applied Biosystems, Foster City, CA, USA) as per QIS 19977. Reaction setup was performed on the MultiPROBE® II PLUS HT EX (PerkinElmer) pre-PCR platform.

5.4 PCR amplification and fragment analysis

DNA extracts were amplified using the AmpFtSTR® Profiler Plus® kit (Applied Biosystems Foster City, CA, USA) as per QIS 19976. Reaction setup was performed on the MultiPROBE® II PLUS HT EX (PerkinElmer) pre-PCR platform.

5.5 Capillary electrophoresis and fragment analysis

PCR product was prepared for capillary electrophoresis using the manual 9+1 protocol (refer to Project 15 and QIS 19978). Capillary electrophoresis was performed on an ABI Prism® 3100 Genetic Analyzer (Applied Biosystems, Foster City, CA, USA) under the following conditions: 3kV injection voltage,10 sec injection time, 15kV run voltage,100µA run current, and 45min run time. Data Collection Software version 1.1 was used to collect.



Rage 12 of 34

raw data from the ABI Prism® 3100 Genetic Analyzer. Fragment size analysis was performed using GeneScan 3.7. Allele designation was performed using Genotyper 3.7, with thresholds for heterozygous and homozygous peaks at 150 and 300 RFU respectively The allelic imbalance threshold is 70%.

Results and Discussion

Criteria for acceptance

Various commercial DNA extraction kits (as per Table 1) were evaluated in order to compare their performance against the current in-house Chelex[®] protocol. These kits were chosen because they were designed specifically for forensic samples and representative of the DNA capture technologies that were out on the market. Furthermore, these kits were manufactured by leaders in the field of DNA extraction technologies with a track record performance in supplying the forensic market with new and reliable products.

We assessed both magnetic bead and silica-based membrane technologies as the automated MultiPROBE® II platforms on which these systems will ultimately be operating on are fully compatible with both systems. The criteria against which the different kits were assessed on include:

- 1. Total DNA yield; the kit must yield sufficient DNA to perform multiple downstream tests such as DNA quantification and PCR amplification.
- Quality of the resulting DNA profiles; the kit should be able to isolate DNA of a suitable quality for PCR amplification of STR loci, in order to generate DNA profiles that are suitable for forensic and human identification purposes.
- Ability to remove inhibitors; the kit must be able to remove common inhibitors present in mock forensic samples (e.g. hemoglobin) using the basic manufacturer's procedure without the use of organic solvents.
- Usability; the kit (and the manufacturer's recommended protocol) must be user. friendly. The necessary steps to prevent cross-contamination should also be described in the protocol. The extraction process should be able to be completed, in a reasonable amount of time, comparable to the current procedure.
- Availability of validated forensic protocols; the kit, including the manufacturer's protocol, must be validated for forensic use, either by the manufacturer or by a forensic laboratory, as determined from statements in the manufacturer's protocol or availability of publications in peer-reviewed journals.

 Availability of a validated MultiPROBE® If PLUS test file; the kit should have a second
- validated MPT file for use on the MultiPROBE® II PLUS HT EX platform.

Assessment of points 1, 2 and 3 was performed through experimentation. Point 4 was assessed based on operator feedback. This report provides results for points 1, 2, 3 and A more extensive assessment of Point 3 was performed on the kit that was found to provide the best results for points 1, 2, 3 and 4 and is reported in Project 11. For points 54 and 6, the availability of validated protocols for all kits evaluated is outlined in Table 3.

The acceptance criteria were strictly adhered to in order to objectively evaluate the different systems. Out of all five DNA extraction technologies, there only existed a validated MultiPROBE® II PLUS test file for the DNA IQ™ system (Table 3). Although this was considered an advantage for DNA IQ™, we did not prematurely dismiss any of the other kits prior to evaluation. We decided that if a kit significantly outperformed the rest, and did not have a validated MPT file already created, that we would create a novel program file



with the kit manufacturer's assistance. This, however, would only be decided at the conclusion of the evaluation process.

Table 3. An assessment of available validated protocols for the various kits that were evaluated by Forensic Biology FSS.

Kit	Availability of va	Availability of validated
DNA IQTM	. 1	
QIAamp® DNA Micro	✓	and the second
ChargeSwitch®	🗸	W K
forensicGEM TM		The state of the s
NucleoSpin® 8 Trace	·	

The results and discussion for each of the kits that were evaluated, in comparison to Chelex®, are provided in the following sections. Refer to Tables 4 and 5 for quantitation results for cell and blood samples respectively. Yield calculations for Chelex®, samples assume a final elution volume of 150µL.

6.2 Evaluation of DNA IQ™

The DNA IQ[™] system uses a novel paramagnetic resin for DNA isolation of consist of two steps: (1) lysis of the biological material on solid support; (2) using the paramagnetic resin to bind DNA, which allows washing of the resin-DNA complex while the resin is immobilised by a magnetic force, in order to remove the lysis reagent and inhibitors in solution.

The manufacturer's method required the use of the MagneSphere® Magnetic Separation Stand. This magnetic stand is used for the separation of the magnetic pelletin 12 samples at a time. The time to process a batch of 12 samples using the DNA IQ™ system takes about 3 hours, including 30 minutes of incubation time.

Three controls were run with each extraction batch: (1) a negative extraction control (empty tube); (2) a positive extraction control (QC dot saliva or blood depending on the extraction) and (3) a substrate blank (the substrate with only saline).

Samples were extracted using the DNA IQ™ method as described in the Methods section and eluted using 100µL Elution Buffer. Due to volume loss during pipetting, the final elution volume is actually around 95µL. The same set of samples was also extracted using the inhouse Chelex® protocol for comparison. Tables 4 and 5 display the DNA concentration (ng/µL) and yield (ng) for all cell and blood samples, compared to the results generated by Chelex®.



age 14 of 34

Table 4. Quantitation values for cell samples on different substrates after extraction by Chelex® and the evaluated DNA extraction kits,

Cette samples	Chalex			DNA	DNA IQ QLAamp DNA Micro			. Charge	Switch.	forunalcGEM		Wicked Sale form	
		Concentration	Yleid*	Concentration	YTeld	Concentration	Yfeld -	Concentration	Ylald	Concentration	Yield	Concentration	Wield
Sample ID	Substrate type	ng/ut.	ng	ng/uL	ng	ng/uL	, ng	ng/uL	ng	ng/ul.	ng	T no.ul	ng
5 (4uL, neat)	FTA	0.058800	11,877600	0,028700	2.670009	0.006030	0.271350	0.023900	3.585000	0.025700	2.821400-	4 80 018200 -	2000 t
SS (cotton) (4uL, neal)	Cotton sweb	0.007410	1.111500	0.098000	9,800000	0.025800	1.151000	0.096700	14.505000	0.083300	18.825500	5 7 0 088900	6.89000
C5 (4uL, nest)	Cotton cloth	0.001480	0,222000	0.050700	5.070000	0.004880	0.219600	0.014900	2.235000	0.037400	7-554800)	0.071900	7.19000
05 (4uL_ neat)	Danim doth	0.062360	0.354000	0.028200	2,820000	0.002180	0.097200	0.003250	0.487500	0.041300	28008	0.043900	4.39008
R14_Neat 4uL	Rayon swab	0.001620	0.243000	0.018000	1,000000	0.000000	0.000000	0,011800	1.770000	0.024000	848000	0.031600	3.18000
R15_Neat 4uL	Rayon swab	0.001580	0,237000	0.019400	1.940000	0.005050	0.227250	0.018100	2,715000	0.019003	3.838000	0.115000	:1,5000
R16_Neat 4uL	Rayon swab	0.000000	0.000000	0.015500	1.550000	0.005610	0.297450	0.027400	4.140000	0.010300	82800	0.057400	5.74000
R17 Neut Aut	Rayon swab	0,000000	0.000000	0.011200	1.120000	0.007310	0,328950	0.005910	0.885500	0.019700	1 miles	20,029900	2,99000
Rayon maan (Nast)		0,000800	0.120300	0.014025	1,402500	0.004743	0.213413	0.015803	2.370376	0.018500	3.737000	10.058525	á.85250
Rayon STD (Neat)		0.000924	0.138586	0,004291	0,429137	0.003300	0.148430	0.009195	1.379299	20.0G5285	1,087483	20.039683	3.96833
4 (4uL, 1/4 dilution)	FTA	0.010300	2.080600	0.005790	0,579000	0.005270	0.237150	0.001250	0.189000	0.007510	0.786020	0.005710	0.57100
64 (ootton) (4uL, 1/4 dilution)	Cotton swab	0.000756	0.113400	0,019000	1,9000000	0.001460	0.066600	0.031600	4.740000	0.030900	5.291800	0.000500	■.95000
24 (4uL, 1/4 dilution)-	Colton doth	0,000541	0.081150	0.015200	1,520000	0.040900	1.840500	0.000000	0.0000000	0,011600	2.3432000	0.018900	1.89000
04 (4ut., 1/4 düution)	Davim doth	0.000000	0.000000	0.045800	4,580000	0.041800	1,881000	0.001720	0.258000	0.013400	2.706800	3,0,017800	-,78000
R10 1/4 4uL	Rayon awab	0.000558	0.083700	0.005740	0,574000	0.001600	0,681000	0.002860	0.429000	0.002950	0,5959003	0.005750	0.67500
R(1_1/4 4ul.	Rayon swab	0.000000	0.000000	0,002560	0.256000	0.061300	0.058500	0.005150	0.922500	0,002020	0.408040	0.001220	0.12200
112_1/4 4UL	Rayon swab	0.000898	0.134700	0.009750	0.975000	0.005570	0.250650	0.006560	0.984000	20,002343	0.472680	0.010200	02000
113 1/4 4uL	Rayon swab	0.000433	0.084950	0,000000	0.000000	0.001550	0.089750	0.001350	0.202500	0,004030	0,814060	9.015000	1,60000
Rayon mean (1/4)		0.000472	0.070838	0.004513	0.451250	0.002555	0.114975	0.004230	0,634500	0.002835	0.572670	0.009545	C.85450
Rayon STD (1/4)		0.060371	0.055667	0.004208	0.420766	0.002020	0.090915	0.002536	0,380328	0.000865	0,178801	0.005198	€81958
3 (4ut., 1/8 dilution)	FTA	0.008170	1.650340	0.008410	0.641000	0.000000	0.000000	0.000000	0.0000000	0.0083 (24.2	0,643620	0.000000	C.00000
33 (cotton) (4uL, 1/8 dilution).	Cotton swab	0.003710	0.556500	0.012190	1.210600	0.001680	0.075600	0.009130	1.369500	0.0000000000000000000000000000000000000	0,601940	0.014900	1.49000
3 (4ut., 1/8 dilution)	Cotton cloth	0.002600	0.390000	0.010400	1,040000	0,000000	0.000000	0.000355	0.053260	0.0000112001	21 12020	0.006570	C,65700
3 4uL, 1/8 dikidon)	Danim doth	0.000739	0.110850	0.007630	0.763000	0.015100	0.679500	0,000000	0.000000	0.007710	1501540	0.000000	0,00000
R6 1/8 4uiL	Rayon sweb	0.000000	0.000001.0	0.001010	0,101000	0,000000	0.000000	0.000597	0.104550	0.003,00	0.6262 W	0.007860	0.76800
R7 1/8 4ul.	Flayon swab	0.000000	0.000000	0.000982	0.098200	0.000000	0.000000	0.00000.0	0,000000	0.003160	0.638320	0.013800	1.38000
R8_1/8 4uL	Rayon swab	0.000000	0.000000	0.001540	0.154000	0.900000	0.000000	0.003390	0,508500	0.000000	0.0000003		
R9 1/8 4UL	Rayon awab	0.000739	0.110850	0.003050	0,305000	0.000000	0.000000	0.003350	0.504000	0.000000	0.000000		7.4
Rayon mean (1/8)		0.000185	0.027713	0.001846	0.164550	0.000000	0.000000	0.001862	0.279283	0.007565	0.316130	0,010830	1.08300
Rayon STD (1/8)		0.000370	0.035425	0.000971	0.097088	0.000000	0.000000	0.001770	0,765562	0.001807	0.385009.	0.004200	2,42002
2 (eut., 1/18 diuton)	FTA	0.000000	0.000000	0.000935	0.093500	0.003940	0.177300	0.000000	0,000000	0.001840	_00187650%	0.000000	0 00000
52 (cotton) 4uL, 1/15 dilution)	Cotion swab	0.000000	0.000000	0,002900	0,280000	0.000000	0.000000	0.001520	9.228000	0.002280	0,450560	0.0000000	0.00000
C2 (4ML, 1/16 dilution)	Cotton data	0.000000	0.000000	0.005010	0,501000	0.001870	0.084150	0.000000	0.0000000	0.000741073	0,149582	0.000000	0.00000
02 (4uL, 1/18 diktion)	Denlm data	0.000000	0.000000	0.002870	0,287000	0.000227	0.102150	0.000000	0.000000	0,0000000	\$0.000000	0,000000	0.00000
R2 1/16 4uL	Rayon swab	0.000000	0.000000	9.000717	0.071700	0,000000	0.000000	0.000000	9.000000	0.061200	0.242400	0.000000	0.0000.0
73 1/16 4uL	Rayon awab	0.000000	0.000000	0.000000	0.000000	0.000000	0.000000	0.000000	0.000000	0.000632	0.127664	3.0.0022803	0.27800
R4_1/16 4uL	Rayon swab	0.000720	0.108000	0.002230	0.223000	0.000000	0.000000	0.003640	0.548000	0.002590	0.523180	0.004480	0.44800
R.5_1/16 4uL	Rayon swab	0.000000	0.000000	0.000000	0.000000	0,000000	0.000000	0.000000	0,002000	0,000788	0.159176		9
Rayon mean (1/16)	,-11 01100	0.000180	0.027000	0,000737	0.073675	0.000000	0.000000	0.000910	0.136500	0.001303	0.253105	70 302253	0.22533
Rayon STD (1/16)		0.000760	0.054000	0.001051	0.105131	0.000000	0,000000	0.001820	0,273000	0.000891	0.180012	0.002240	March 125 No.

Page 15 of 34

Table 5. Quantitation values for blood samples on rayon swab substrates after extraction by Chelex® and the evaluated DNA extraction kits.

Blood samples	Chel	ex.	· DNA	IQ	QlAamp Di	NA Micro	ChargeS	witch	forens	ICGEM	NucleoSpir	1 8 Trace
	Concentration	Yeald*	Concentration	Yleld	Concentration	Yield	Concentration	Yield	Concentration	Yleid	Concentration	Yield
· · Sample ID	ng/uL	ng	ng/uL	ng	ng/uL	ng	ng/ul.	ng .	ng/ut	.09	aring/ut	ng
R14 (Neat)	2.37	355,5	0.482	48.2	2.31	103.95	0.751	112.65	10,00833	68266	1.16	116
R15. (Neat)	1.42	213	0.078	7.8	. 3,58	161.1	0.754	113,1	20.0066	1.3332	2.61	261 .
R16 (Neat)	0.512	. 76,8	0.356	35.6	3.32	149.4	0.929	139.35	0.0046	.0.9292	1.61	161
R17 (Neat)	0.934	140.1	0.467	46.7	2.46	110.7	0.916	137 4	0.00727	1.46854	2.18	218
Mean (Neat)	1.3090	196.3500	0.3458	34.5750	2.9175	131.2875	0.8375	125.6250	0.0067	1 3534	1.8900	189.0000
STD (Neat)	0.7987	119.8085	0.1871	18.7137	0.6270	28.2137	0.0983	1497451	0.0016	03173	0.6361	63.6082
R10 (1/4)	0.219	32.85	0.238	23.8	0.227	10.215	0.219	32 85	0.002/11	0.42622	0.611	61.1
R11 (1/4)	0.0845	12,675	0.198	19.8	1.72	77.4	0.101	15:15	0.000597		0.3	30
R12 (1/4)	0.216	32.4	0.195	19.5	4.59	206.55	0.0673	10:095	0.00128	0.25856	0.251	25.1
R13 (1/4)	0.165	24.75	0,136	13.6	0.657	29.565	0.0787	14.805	==10.000166U	0.33532	0.227	22.7
Mean (1/4)	0.1711	25.6688	0.1918	19.1750	1.7985	80.9325	0.1165	17/4750	0.0014	0.2852	0.3473	34.7250
STD (1/4)	0.0628	9,4262	0.0420	4.2019	1.9539	88.3776	0.0698	10.4628	0.0006	0.1294	0.1784	17.8438
R6 (1/8)	6,88	1032	0.0554	5.54	0.0936	4.212	0.094	14.1	0.0126	2.5452	0.154	15.4
R7 (1/8)	0.164.	24.6	0.114	11.4	0.175	7.875	0.0735	11.025	0.00174	0.35148	0.148	14.8
R8 (1/8)	0.286	42.9	0.145	14.5	0.123	5,535	0.0521	7.815	0.00363	0.73326	0.178	17.8 .
R9 (1/8)	0.513	76.95	.0.125	12.5	0.0151	0.6795	0.0939	14.085	0.00167	0.33734	0.0819	8.19
Maan (1/8)	1.9608	294.1125	0.1099	10.9850	0.1017	4.5754	0.0784	11.7563	0.00490	0.9918	0.1405	14.0475
STD (1/8)	3.2827	492,4030	0.0385	3,8501	0.0668	3.0066	0.0200	2.9991	0.0052	1:0517	0.0411	4.1145
R2 (1/16)	0.0405	6.075	0.0792	7.92	0.0349	1.5705	0.0347	5.205	0.00757	1.52914	0.0766	7.66
R3 (1/16)	0.0104	1.56	0.0566	5.66	0.0454	2.043	0.027	4.05	0.00667	34734	0.0923	9.23
R4 (1/16)	0.0337	5.055	0.0847	8.47	0.0386	1.737	0.0197	2.955	0.00544	1.09888	0.0588	5.88
R5 (1/16)	0.0323	4.845	0.109	10.9	0.0276	1.242	0.021	3.15	-0:00245	0.4949	0.874	87.4
Mean (1/16)	0.0292	4.3838	0.0824	8,2375	0.0366	1.6481	0.0256	3.8400	0.0055	1 (176	0.2754	27.5425
STD (1/16)	0.0131	1.9577	0.0215	2.1515	0.0074	0.3341	0.0068	1.0274	30,0022	0.4510	0.3993	39.9285

Page 16 of 34

Comparison of quantitation results for cell samples

Refer to Table 4 for observed data. Using DNA IQ™, neat cell samples displayed higher quantitation results for both cotton and rayon swabs, and also for cotton and denim cloth materials. Only for the FTA® card was the result higher for the Chelex® sample. For 1/4 dilutions, DNA IQ™ results were higher than Chelex® results. For 1/8 dilutions, both protocols showed similar results for most sample types. Rayon swabs produced zero quantitation values for Chelex®, but exhibited consistent results for DNA IQ™ Fold 1/6 dilutions, most Chelex® samples were undetermined, whereas most DNA IQ™ samples yielded quantitation results.

Only three dilution samples extracted by DNA IQ™ gave zero quantitation values. In contrast, fourteen Chelex® samples gave zero quantitation results. This suggests that the DNA IQ™ sample recovery rate is 111% greater than that of the Chelex® protocol for cell samples.

Comparison of quantitation results for blood samples

Refer to Table 5 for observed data. For this experiment, only rayon samples (in quadruplicate) were tested. Neat blood samples showed higher concentration, esuits when extracted using Chelex®. The 1/4 dilutions showed similar results for both methods. The 1/8 dilutions showed better results for Chelex®, but this was primarily due to an outlier result for one of the replicates (highlighted red in Table 4) that resulted in a concentration value 1300% greater than the remaining samples. This occurrence could be the result of inaccurate pipetting during mock sample creation or variability in the Chelex® method specifically the inconsistent final elution volumes. For the 1/16 dilutions, the DNA IQ™ results were better. All DNA IQ™ results were more consistent and reproducible than Chelex® results.

Overall, samples that were extracted using DNA IQ™ showed quantitation results that were similar to or better than samples that were extracted using Chelex®. For cell substrates, 44% of Chelex® samples gave zero quantitation results, compared to only 9% for DNA IQ™ samples. All blood substrates generated quantitation results that were similar for both methods. Furthermore, DNA IQ™ generated results that were more sensitive, consistent and reproducible across multiple replicates.

Comparison of DNA profiles

Cell samples that were extracted using the DNA IQ™ method gave DNA profiles with more alleles compared to extractions performed using Chelex® (Table 6). Overall, DNA IQ™ resulted in 282 reportable alleles (excluding Amelogenin), compared to 89 alleles resolved by Chelex®, or in other words samples extracted using DNA IQ™ generated 216% more reportable alleles compared to samples extracted using DNA IQ™ generated 216% more reportable alleles compared to samples extracted using Chelex®. For neat cell substrates SNA IQ™ samples generated full profiles in all instances except 2: an X,X+14 for the FTA substrate and an X,X+16 for a rayon swab replicate. All rayon samples extracted by Chelex® did not produce any profiles at all, in contrast to the full profile results using DNA IQ™. DNA IQ™ also gave more reportable alleles for the lower dilutions compared to Chelex®. Additionally, DNA IQ™ was able to yield full profiles from denim substrates, compared to Chelex® which yielded no profiles at all. This observation indicates the superiority of the DNA IQ™ system for removing and overcoming inhibition due to denim dye. Only one occurrence of allelic imbalance (68% at D13S317) was encountered in all 64 samples.



Rage 17 of 34

Table 6. Comparison of DNA profiles for cell substrate samples extracted using either Chelex® or DNA iQ™.

CELLS		Method: Ch	relex				
Ollution	FTA	Cotton	Rayon sw	wbs	Cottón	Denim	
	Profits	Profile	Sample#	Profile	Profile	Profile	
Neal	XX+18	XX+18	R14	NSD	X,X+8	NRVISD	
-		与 学生是多级	R16	NR/NSD	10 miles		
	P 18575	प्रदे श होते हुए	R16	NSD	130	100	
,	200	159112	R17	NSD		- 101	
Dil 1/4	X,X+18 -		R10	NSD	NRM .	NRVNSD	
			R11	NSD	272600	100	
		164	R12	NSO	2000	20000000000000000000000000000000000000	
		A TOTAL CONT	R13	NSD	2007	27.00	
DE 1/8	XX+17	XX+3	R6	NSD	X,NR+3	NR/NSD	
			R7	NSD	2.5		
			R8	NSD	THE REAL PROPERTY.		
		The Course	R9	NSD		E 的 EN	
Dil 1/18	NSD .	NSD	R2	NSD	NSD	NSD	
	distribution of	75/70	R3	NSD	COLD STATE	100	
	100000000000000000000000000000000000000	Street Street	R4	NSO	Part Control		
				-	CONTRACTOR OF STREET, S. S.	THE RESERVE TO SERVE	

Dilution	FTA	Cotton swebs	Rayon swab	s	Coffon	Dentmiss	
	Profile	Profile	Sample#	Profile	Profile	Profile (S. D.	
Neat	X,X+14	X,X+18	R14	XX+18.25%	XX+18	XX+18	
	THE REAL PROPERTY.		R16 ×	AI@D13(68%	在水水流流		
	THE STREET	21 TO 1	R18 3	XX+18 17	A STATE OF		
	200		R17 .公常認	X;X+18	1000	200	
DH 1/4	X,X+17	X,X+18	R10 375	XNR+3	X,X+16	XX+18	
	E TANK	性的主流的	R11	NR/NSD:	4.52	The Second	
			R125	X X+6.	1	125	
	經過數學	76 F	8.13	NRNSD		42.46.29	
Dil 1/8	XX+8	X,X+18 #	Bd	NR/NSD	X,X+17	X,X+17	
		100	RICE	NR/NSD	200		
		CERTIFICATION OF THE PERSON OF	Ru	NRMSD	E CAN		
	200		Re.	NR/NSD			
DII 1/16	NSD	XX+4 1833	R2	NR/NSD	NR/NSD '	NRINSD	
	100 min		ĖJ	NSD .			
		第二条	Rij	NR/NSD		G 6400 (2)	
	然如我就是		FLB	NSD 38			

For blood samples, only rayon substrates were extracted using the DNA IQ." System as these were deemed sufficient for observing the effects of heme inhibition (without the need to factor variable substrate types). Almost all samples generated full profiles of a sufficient number of reportable alleles for matching purposes (Table 7). For neat samples extracted by Chelex®, no profiles were resulted from the FTA®, cotton swab or denim samples, indicating possible heme inhibition that could not be removed by the Chelex® protocol For rayon samples, 19% of those extracted by Chelex® did not generate a profile, whereas DNA IQ™ yielded full profiles for all dilutions except two neat samples. Reworks of the two failed samples were performed but yielded the same NSD results. These failed results appear to be outliers, as all other dilutions yielded the expected results. It was observed that results from blood samples on rayon swabs were more likely (32%) to exhibit allelic imbalance at Amelogenin when extracted using the DNA IQ™ system.

Table 7. Comparison of DNA profiles for blood substrate samples extracted using either Chelex or DNA IQ™

BLOOD		Mediod. Cir	DIGA					Maniou. DRA IG		1
Dilution	FTA	Cotton swabs	Rayon sw	rabs	Catton	Denim		Rayon swabs		-
	Profile	Profile	\$ample#	Profile	Profile	Profile		Sample#	Profile	
1000000		NSD	R14	X,Y+18	X,Y+18	NSD		R14	NSD	į
			R15	X,Y+18		100 Kar 123		R15	NSD	ľ
	高级 电路线		R16	X,Y+18		0.6 92		R16	X.Y+18(AL@AMEL)	i
	- The Part of the		R17	NR/NSD		建筑直线		R17	X,Y+18(AI@AMEL)	į
1000		X,Y+15	R10	Not Uploaded	X,Y+18	X,Y+18		R10	X,Y+18	1
	建	heer-off	R11	X,Y+18		建筑建筑	2 16 7 5 6 6	R11	X,Y+18(Al@AMEL)	b.
		10,270	R12	X,Y+18				R12	XY+18	3
		#T07%/2	R13	X,Y+18	建筑域数	2004		R13	X,Y+18	1
Dil 1/8 X,Y+1	X,Y+18(A!@I	XY+18	R6	X,Y+18		X,Y+18		R6	X,Y+18(Al@AMEL)	ı
	Service Control		R7	X,Y+18				R7	X,Y+18	
			R8	NRVNSD				FLB	X,Y+18	
			R0	X,Y+18	1000	200		R9	XY+18	0
	X,Y+18	X,Y+18	R2	X,Y+18	X,Y+18	X,Y+18	-	R2	XY+18(AI@AMEL)	17
	RECEIVED.	(4.6 ± 1)		X,Y+18	No.		200	R3	XY+18	
		45-14	R4	X,Y+18				R4	XY+18	1
	14-14-15 AV AV AV AV	of the transfer over the latest and the		_	The second second second	The second second				10



Page 18 of 34

We found the DNA IQTM system yielded results that were either comparable or better than results generated by samples extracted using the in-house Chelex® protocol, both in terms of quantitation values and DNA profile quality and completeness.

6.3 Evaluation of QlAamp® DNA Micro

The QIAamp® DNA Micro kit was designed for the purification of genomic and init school and DNA from small sample volumes or sizes, as often encountered in forensics. The system uses a silica-based membrane to accommodate DNA binding and purification using special buffers, followed by elution in buffer or water, resulting in purified DNA that is free of proteins, nucleases and other impurities.

The QIAamp® DNA Micro system consists of four steps: lysing, binding, washing; followed by elution:

- Lysis Small samples are lysed under highly denaturing conditions at elevated temperatures under the presence of Proteinase K.
- Binding Using Buffer AL and ethanol, DNA is adsorbed into the silica-gel. 7
 membrane of the column by centrifugation or application of awacum. The buffer is
 formulated so that proteins and other components are not retained in the
 membrane.
- Washing While DNA is bound to the silica membrane, contaminants are efficiently washed away using a combination of two wash buffers
- Elution DNA is eluted in a small volume of Buffer AE or sterile water, yielding concentrated DNA.

The QIAamp® protocol involves 5 tube transfers and therefore takes approximately 5 hours to perform a manual extraction of 12 samples. The same set of samples that were used for the DNA IQ™ evaluation was also used to evaluate QIAamp® DNA Micro. Each extraction batch included a positive and negative control, and also a substrate blank. DNA,was eluted in 45µL volume.

Comparison of quantitation results for cell samples

Refer to Table 4 for observed data. Twelve samples extracted by QIAamp® gave Zero quantitation values, compared to fourteen samples by Chelex®. Despite the low elution volume of 45µL in the QIAamp® protocol that serves to concentrate the purified DNA, quantitation results for all samples were comparable for both DNA extraction methods.

Comparison of quantitation results for blood samples

Refer to Table 5 for observed data. Blood on rayon swab samples displayed wide variation between replicates. For neat samples, the total yield is comparable to Chelex®, however lower dilutions (1/8 – 1/16) suffer from inconsistencies. One of the 1/4 dilution replicates displayed an unexpectedly high quantitation value that was more than 3x greater than the Chelex® average yield, but this can be attributed to inaccurate pipetting, or pipetting of a non-uniform sample mixture, during mock sample creation.

A possible reason as to why the quantitation results for both cell and blood samples were inconsistent is because the QIAamp® DNA Micro protocol uses five sets of collection tubes for supernatant transfer, therefore possibly causing sample lost during multiple sample transfers from one tube to another.



19 of 34

Comparison of DNA profiles

Cell samples that were extracted using the QIAamp® protocol showed profile results that were either comparable or worse than samples that were extracted using the Chelex® protocol (Table 8). Out of 32 samples, only one QIAamp® sample resulted in a full profile (X,X+18). QIAamp® samples failed to produce full profiles for all but one (n = 8) of the neat samples. Overall, QIAamp® resulted in 86 reportable alleles compared to 89 alleless resolved by Chelex®. Some of the QIAamp® allele calls are inconsistent, e.g. the result for 1/4 dilution on cotton cloth was slightly better than the neat sample. This is further exemplified by the denim substrate samples. The QIAamp® method did not appear to effectively overcome inhibition caused by the denim dye as observed from the resulting profiles.

Table 8. Comparison of DNA profiles for cell substrate samples extracted using either Chelex, or QIAamp DNA Micro.

CELLS

Mathod: Chels

Dilution	FTA	SWabs	Rayon sw	ebs	Cotton	Donlm
	Profile ·	Profile	Sample#	Profile	Profile	Profile
Nest	XX+18	X,X+18	R14	NSD	X,X+8	NR/NSD
	CONTRACTOR OF THE PARTY OF THE		R15	NRANSD		
	SERVICE SERVICE	465 (25.15)	R16	NSD		化的图像
	SEE CANADA	a decided a	R17	NSD	200	4.600
DII 1/4 .:	XX+18 .	XNR+3	R10	NSD	NR41	NR/NSD
	成的似乎更是原		R11	NSD	1000	
			R12	NSD		化 限30
	Section 15	THE RESERVE	R13	NSD		拉拉拉
DII 1/8	XX+17	X,X+3	R8	NSD	XNR+3	NRINSD
			R7	NSD .		
			R8	NSD	医皮脂的	"大大"
	and the second second second	ALC: NO PERSONAL PROPERTY.	R9	NSD	T. Maria Sale	(44.44)
QII 1/18	NSO	NSD .	R2	NSD	NSD .	NSD .
			R3	NSD		
			R4	NSD.	ENDOUGH A	
	基础的时间代的结构	建设水位。	R6	NSD		

CELLS Mothod: Qlanno DNA Micro

Diludon	FTA	Cottop:	Reyon swa	bs som	Cotton	Denim
	Profile	Profile 12	Simple# .5	Profito	Profile	Profile
Neat	X,X+12	X,X+18	原理 点型	NSO	NR,NR+2	NRINSD
	10000000	300000	数15.5%。200	NRINSD	250	
	1	面原物理	RIGHT	XNR+6		
		16/17/19	R17.题能改	NR/NSD	1000	95 WH W
DII 1/4	XX+14	XX+15 .	R1024	NSD.	NR,NR+3	NRINSD
	A CONTRACTOR	(150 PM)	RUSS	NSD	THE REAL PROPERTY.	15 A SW
		6.5	R1238	NE NE L	COLUMN TO SERVICE	经外发票件
		E MANAGEMENT	R(186	NR/NSO		SHOW SHOW
Dii 1/8	NSD+2	XNR+6	RBIX	NSD.	NR/NSO	XNR+7
		100	RANGE	NSDS 18		经规则
	A STATE OF THE REAL PROPERTY.	5.50	RESEL	NSD		
	- A - A - A - A - A - A - A - A - A - A	100 A	ROTTO:	NSD		100 23000
D) 1/18	NR/NSD	NRVNSD	RZ类体生	NSD	NSO: " P	NSO .
	100 C	2000年1月1日	R3 1000	NSD/SON		the street of
			R4 TERRES	NSDE		
	the Land	1	R5 PACK	NSD		

Table 9. Comparison of DNA profiles for blood substrate samples extracted using either Chales of QIAamp® DNA Micro.

BLCO

fethod; Chelex

Method: QIAamp DNA Micro

Dilution	FTA	Cotton	Rayon sw	abs	Cotton	Denim		Rayon swa	bs
	Profile	Profile	Sample#	Profile	Profile	Profile		Sample#	Profile
Neat	NSD	NSD ·	R14.	X,Y+18	X,Y+18.	NSO	1	R14	X,Y+18
		NAME OF	R15	X,Y+18				R15	XY+18
			R18	X,Y+18				R16	X,Y+18
		E40368E	R17	NR/NSD	The same year	DESCRIPTION OF THE PARTY.		R17	NR,Y+15
011/4	X,Y+18	X,Y+15	R10	Not Uploaded	XY+18	X,Y+18	1	R10	X,Y+18
			R11	X,Y+18		28 (I-00	特温系管系	R11	XY+17
			R12	X,Y+18				R12	X,Y+18
	333000		R13	X,Y+18		54.00		R13	X,Y+18
011 1/8 .	X,Y+18(At@t	X,Y+18-	R6		X,Y+18	X,Y+18		R2	X,Y+18
		No.	R7	X,Y+18				R3	X,Y+18
		1000	R8	NR/NSD		FUTARS	是创意的问	R4.	X.Y+18
	是的可能的		R9	X,Y+18		2727		R5	X,Y+18
NI 1/16		XY+18	R2	X,Y+18	X,Y+18 -	X;Y+18		R2	X,Y+18(AI@D8,D18)
			R3	X,Y+18	7 (F) (F)		550 200	R3	X,Y+18
			R4	X,Y+18				R4	X,Y+18
	建筑公园 园		R5	X,Y+18			16170 School	R5	X,Y+18



age 20 of 34

For blood samples on rayon swabs, 87.5% of QIAamp® samples resulted in full profiles, compared to 81.25% of Chelex® samples (Table 9). Out of all QIAamp® rayon swab samples, only one of the 1/16 replicates displayed allelic imbalance (in D8S1179 and D18S51).

6.4 Evaluation of ChargeSwitch®

The ChargeSwitch® technology (CST) is another magnetic bead-based technology that provides a switchable surface charge dependent on the pH of the surrounding buffer environment to facilitate DNA isolation from small forensic samples. In low pH conditions the ChargeSwitch® beads have a positive charge that allows negatively charged DNA to bind. In this environment, proteins and other contaminants are not bound and can be washed away. By using a low salt elution buffer at pH 8.5, the charge on the bead surfaci is neutralised and DNA can be eluted for Immediate use in downstream forensic applications.

The ChargeSwitch® Elution Buffer (E5) that is supplied with the kit is used to provide an environment with a pH of 8.5 that promotes dissociation of bound DNA from the magnetic beads and therefore efficient elution of purified DNA. However, TE buffer with a pH between 8.5 – 9.0 can also be used for elution. TE buffer outside of this pH range should not be used. The use of water for elution is also not recommended.

The manufacturer's method required the use of the MagnaRack™ two-piece magnetic separation rack that consists of two components: a magnetic base station and removable tube rack. The tube rack holds up to 24 microcentrifuge tubes and fits onto the magnetic base station in two different positions associating the row of 12 neodymlum magnets with a single row of 12 tubes for simple 'on the magnet' and 'off the magnet' processing. The times to process a batch of 12 samples using the ChargeSwitch® system takes about 3.5 hours; including 30 minutes of incubation time. Each extraction batch included a positive and negative control, and also a substrate blank. Purified DNA samples were eluted in 150µL Elution Buffer (E5).

Comparison of quantitation results for cell samples

Refer to Table 4 for observed data. For cells samples, ChargeSwitch® performed moderately better compared to the current in-house Chelex® method. When comparing the quantitation values, ChargeSwitch® produced higher quantitation values for cotton and rayon swabs over all dilutions as well as the neat samples of cotton shirt and denim jeans. For other cell samples, ChargeSwitch® performance was comparable to the Chelex® results.

Comparison of quantitation results for blood samples

Refer to Table 5 for observed data. ChargeSwitch® quantitation results for blood samples on rayon swabs were lower but more consistent than Chelex® results.

Comparison of DNA profiles

Cell samples that were extracted using the ChargeSwitch® system showed profile results that were comparable to samples that were extracted using the Chelex® protocol (Table 10). Overall, ChargeSwitch® resulted in 138 reportable alleles compared to 89 alleles resolved by Chelex®. ChargeSwitch® performance for cell samples on FTA® cards was poor for any samples less than the neat dilution. Profiles for both cotton swab and cotton cloth samples were slightly better for ChargeSwitch®, and results for neat samples on rayon



250 21 of 3

swabs outperformed Chelex[®]. However, the ChargeSwitch[®] system was unable to overcome inhibition in denim samples, and did not yield any DNA profiles at all, despite displaying quantitation results for the neat and 1/4 dilution.

Table 10. Comparison of DNA profiles for cell substrate samples extracted using either Chelex of ChargeSwitch.

Dilution	FTA	Swabs	Rayon sw	abs	Cotton	Denim
	Profile	Profile	Semple#	Profile	Profile	Profile
Neat	XX+18	X,X+18	R14	MSD	X.X+8	NR/NSD
	E STATE OF	经有效不同	R16	NRVNSD		TO THE STATE OF
			R16	NSD	Sec. 35 18.0	all a fall
	自然 。在1886		R17	NSD		Charles Tally
DI 1/4	XX+18 -	X,NR+3	R10	NSD	NR+1	NR/NSD
		Market D.Y.	R11	NSO		Contract of the
			R12	NSD	S. 95 (C. A. A.)	Section 1
	(1) F (2)	1000	R13	NSD		
DI 1/3	- XX+17	X,X+3	R6	NSD	XNR+3	NR/NSD
	ta F50 ta 5		R7	NSD	A SECTION	100
			R8	NSD		V BOSESS
		A CONTRACT	RS	NSD	建筑市场	V 10.70
DII 1/16	NSD	NSD	R2	HSD	NSD	NSD
	12-11-645		R3	NSD		A C (
			R4	NED		WAN 35
	September 1	的 公司的编辑	R6	NSD		Land to

CELLS		Method: Charge Switch							
	FTA	cotton swebs	Reyon swal	15.0	Cotton	Denim Profile			
	Profile	Profile A	Sample#	ProNa	Profile				
Neat	X.X+17	X.X+18	R14	X.X+8	XX+11	NSD			
	CVENT CARE			XX+16	2000	THE PERSON			
		F-1275-C-17	E16 07	X,X+16	在學院的	() X			
	超级数据	100	開釋物學	XX B	20000000000000000000000000000000000000				
DI) 1/4	X,X+9	X,X (18/2)	R10	XXINRIA	NRNR+2	NSD .			
	A STATE OF THE STA	Salary 72	R11	NR/NSD		Code			
		F. 0	R12	X,NR+2	100	2003			
	100	Pa 2 3 4 5	R13	X.NR+NSD	和表表 575				
DH 1/8	NRASD	XX+122522	RB 大大型型的	NSDA	NR/NSD	NSD .			
		CALL SECTION	873AS	NSDO	- A-1	100000			
			File , Sil	WWWSD					
	OF LANDS	思证的推	R9 3	NSD	1000				
DB 1/16	NSD	NR/NSD 微	R23	NR/NSD	NSD	NSD .			
	THE COME	1	FO.San	NSD					
		arakiri Uni Pakir	RASHOE	NSD	SECTION S				
	企业信息	22 54 54	R5 (1935)	NSD	建筑型和	B 50			

For blood samples on rayon swab substrates, all ChargeSwitch® samples consistently yielded full profiles for all dilutions and therefore outperformed Chelex® (Table 11). Two replicates of the lower, 1/16 dilutions displayed allelic imbalance at two different loci: D3S1358 and D7S820, possibly due to stochastic effects that arise from amplifying low concentrations of DNA.

Table 11. Comparison of DNA profiles for blood substrate samples extracted using either Chelex or ChargeSwitch.

BLOOD		Method: Chalex						Method: ChargeSwitch	
Dilution	FTA	Cotton			Cotton	Denim	Rayon swabs		
	Profile	le Profile	Sample#	Profile	Profile	Profile	Sample#	Profile	
Neat	NSD .	NSD	R14	X,Y+18	X,Y+18	NSD	R14	X,Y+18	
			R15	XY+18			R15	X,Y+18	
		7. O. S.	R16	X,Y+18	用数数	100	R16	X,Y+18	
	5/2007		R17	NR/NSD	N 12-21 / 4		R17	X,Y+18	
DII 1/4	X,Y+18	X,Y+15	R10	Not-Uploaded	X,Y+18	X,Y+18	R10	X,Y+18	
	120 W # 20		R11	X,Y+18			R11	X,Y+18	
			R12	X,Y+18	K Z Z Z Z	SUPER	R12	X,Y+18	
		SERVICE STATE	R13	X,Y+18			Rt3	X,Y+18	
DII 1/8	X,Y+18(A1@0	X,Y+18	R6	X,Y+18	X,Y+18	X,Y+18	R6	X,Y+18	
		2.3	R7	X,Y+18	N. Carlot	5.0	R7 .	XY+18	
	AND SECULO		R8	NR/NSD		200	R8	X,Y+18	
	75.274.2	研修通過	R9	X,Y+18			R9	X,Y+18	
DII 1/16		XY+18	R2	X,Y+18	X,Y+18	X,Y+18	R2	X,Y+18	
			R3 .	X,Y+18 .			R3	X,Y+18	
	學者的不够	Charles.	R4	X,Y+18	V-32-5		R4	X,Y+18(Al@D3)	
	22 × 15	自然是10万位	R5	X,Y+18			R5	X,Y+18(AI@D7)	



Page 22 of 34

6.5 Evaluation of forensicGEM™

forensicGEM™ is a novel thermostable proteinase developed as a rapid, cheap and effective single-tube DNA extraction solution for forensic laboratories that was recently released. At the time of testing, the forensicGEM™ system was not yet widely used in field of forensics, however the system has had exposure at various conferences and symposiums, such as the 18th International Symposium on the Forensic Solenges (Fremantie, WA; 2-7 April 2006).

Unlike the other kits that were evaluated, *forensic*GEM™ does not incorporate either magnetic-bead or silica membrane technologies, but instead works on the principle action of a thermostable proteinase in an optimised buffer solution. *forensic*GEM™ is based on the work of Moss *et al.* (2003) who developed the use of EA1 proteinase for the DNA extraction of forensic samples. EA1 proteinase comes from the their ophilic *Bacillus* sp. EA1. EA1 proteinase is Ca²⁺ dependent but is unaffected by a concentration of citrate below 5mM and EDTA below 2mM (Moss *et al.* 2003). For EDTA-stabilized blood, the buffer needs to be supplemented to a final concentration of 200µM ⟨acl₂, Heating a sample at 75°C in the presence of *forensic*GEM™ buffer and *forensic*GEM™ ityses the sample and the proteinase hydrolyses nucleases. At 95°C the proteinase is near inactivated so that an active form will not be carried over into PCR where it would degrade *Tag* DNA polymerase.

The time to process a batch of 12 samples using the *forensic*GEM[™] system takes about 1.5 hours. Each extraction batch included a positive and negative control, and also a substrate blank. The final volume was 100µL for FTA[®] samples and 200µL for all other samples.

Comparison of quantitation results for cell samples

Refer to Table 4 for observed data. For cells samples, *forensic*GEM™ produced aigner quantitation results compared to Chelex® across all dilutions. *forensic*GEM™ also generated the highest yield for all samples, including the 1/16 dilutions. *forensic*GEM™ yielded quantitation results for denim samples (neat and 1/4 dilutions).

Comparison of quantitation results for blood samples

Refer to Table 5 for observed data. *forensic*GEMTM performed very poorly for blood samples on rayon swabs, resulting in the lowest observed yield across all kits that were evaluated. The average yield for all four neat replicates processed using *forensic*GEMTM was 0.6% of the average yield for all Chelex® replicates. The best average yield results were observed for 1/16 dilution samples, where the average *forensic*GEMTM yield was around 25% that of Chelex®. This suggests that the *forensic*GEMTM system is prone to heme inhibition if a neat sample is processed, but can slightly overcome the inhibitory effect if the blood sample is diluted prior to extraction.

Comparison of DNA profiles

forensicGEM™ resulted in 209 reportable alleles for cell samples compared to 89 alleles resulting from Chelex® extracts (Table 12). forensicGEM™ was able to overcome inhibition in denim samples, producing full profiles (X,X+18) for neat and 1/4 dilutions, accurately reflecting the quantitation results. A partial profile (X,NR+7) was obtained for the 1/8 dilution on denim. forensicGEM™ results were also superior than Chelex® for cells on cotton swab down to the 1/8 dilution, but FTA® results were considerably poor.



1Page 23 of 34

Table 12, Comparison of DNA profiles for cell substrate samples extracted using either Chelex[®] or forensicGEM[®].

Dilution	FTA	Cotton swabs	Reyon swabs		Gotton	Denim
	Profile .	Profile	Sample#	Profile	Profile	Profile
Noàt	X,X+18	X,X+18	R14	NSD	XX+8	NRAISD
	No. of Contract of		R16	NR/NSD	14 SEC. 1	
	or that		R16	NSD	4	THE CASE
	9747	(1) (1) (1) (1)	R17	NSD	2000年	1912 (129)
DI 1/4	X,X+18 .	XNR+3	R10	NSD	NR+1	NR/NSD .
	The state of the s		R11	NSD	25.00000	100
	全国的政党	推销 数200	R12	NSD	正常有些性	AREL AN
	不可能的	GROSS CO	R13	NSD	MARKET	
D1 1/8	XX+17	XX+3	R8 .	NSD	XNR+3	NR/NSD
	CHE THE	100	R7	NSD		
	T 10 10 10 10 10 10 10 10 10 10 10 10 10		RB	NSD		10789298
-	10 to	THE STATE OF	R9	NSD	STATE OF THE	

CELLO		Wenton: lou	PRHECIEM	31.	A	10 m
	FTA	Gotton swabs	Rayon swat		Cotton	Denim
	Profile	Profile	Sample#	Profile Park	Profile City	Prome.
Neet	X.X+15	XX+18 ·	R14 ,	XXXII	XX-18	XX+18 *****
	泰田和松松	14.5	R15	XIX+13		
	at the form		R16	XX+3	100	3.72
		### IN S	RIVERS	X.X+ 18		
DII 1/4	X,NR+3	X,X+18	A10SESSES	X.NR THRINS	XX+18	XX+18
	70.00		图11、通路区	NR/NSD)		14.
		A 100		NRINSDA	STATE OF THE PARTY	(A) (A) (A)
	A2017 A10		R13	XNRINKINS	STATE OF STREET	
DB 1/8	NSD		R7	NR/NSOE:	XNR+10	X,NR+7
			R8	NSD TE	ACCEPTAGE	
	A STATE OF		R9762	NRINSD		
		is to proper	R6 " PT # P	NEVISO	亚 大学	
OII 1/18	NSD	NEWSD	R6	NSD "STELL	NSD.	NR/NSD .
	建筑建筑建筑	725 F. DW	R4	NSD		The Control of the
		TO THE REAL PROPERTY.	R3	NSDE SE		
	22/25/20/2	45(7)-75	R2位金融器	NR/NSD	1	
		15 E.S	10	25000		

For blood samples on rayon swabs, only the 1/16 dilutions generated profile results (Table 13). This is indicatory of potential inhibition for higher blood sample dilutions as predicted by the quantitation data.

Table 13. Comparison of DNA profiles for blood substrate samples extracted using either Chales of forensic GEM®.

BLGOD		Method: Chalex					Method: forensicGEM		
Dilution	FYA	Gotton	Rayon swabs		Cotton	Denim	Hayon swabs		
	Profile	Profile	Sample#	Profile	Profile	Profile	Sample#	Profile The Profile	
Neal	NSD	NSD	R14	XY+18 ·	XY+18	NSD	R14	NSD W	
		2.5	R15	XY+18			R15	NSD TO	
			R16	X.Y.+18 .	THE REAL PROPERTY.	1200	R16	NSD SSOR	
	STATE OF THE STATE OF	237	R17	NRASD	建设建设		R17	NSD S	
Di) 1/4	XY+18	X.Y+15	R10	Not Uploaded	X,Y+18 .	X,Y+18	R10	NSD "	
		753-155	R11	X,Y+18	2.5		R11	NSD	
		1000	見12	XY+18		100	R12	NSO	
	(1) (2) 4 (2)		R13	X,Y+18	7000000	1000	R13	NSD	
DII 1/8	X,Y+18(Al@0	X,Y+18 .	R6		XY+18	XY+18	R8	NSD	
	Acres 14	STATE OF THE PARTY	R7				R7 ·	NSD	
	阿尔拉拉斯 自由 4		R8	NR/NSD	2	1762 256	Rô	NSD	
	12.7	532543	R9	X,Y+18 ·			R9	NSD	
DK 1/18	X,Y+18		R2	X,Y+18	X,Y+18	X,Y+18	R2	X,Y+15(AISQD13)	
		Te Section		X,Y+18	2500	500	R3	X,NR+3	
	CAN COMPANY	- 50	R4	XY+18		建新疆 物		NR,NR+5	
	20 多用作品		R6	X,Y+18	The state of the s	13.5	R5	NR.NR+2	

6.6 NucleoSpin® 8 Trace

The NucleoSpin® 8 Trace kit is designed for extraction of genomic DNA from forensic samples. Cell lysis is achieved by incubating samples in a solution containing chaotropic ions in the presence of proteinase K at room temperature. Adding isopropanol to the lysate creates the appropriate conditions for binding of DNA to the silica membrane, a process that is reversible and specific to nucleic acids, inhibitors are removed by washing steps using an alcohol-containing buffer. Pure genomic DNA is eluted in a slightly alkaline elution buffer.

The evaluation of this kit was performed with slight alterations in the manual method to incorporate the use of the MultiPROBE® II PLUS PVM vacuum manifold, together with the



NucleoSpin® 8 Trace Starter Set A containing Column Holders A and Dummy Strips to enable use of the vacuum manifold.

The time to process a batch of 12 samples using the NucleoSpin® 8 Trace system takes about 5 hours, including a 3 hour incubation step. Each extraction batch included a positive and negative control, and also a substrate blank. Purified DNA was eluted in a final volume of 100µL.

Comparison of quantitation results for cell samples
Refer to Table 4 for observed data. NucleoSpin® 8 Trace produced greater mean concentration values and mean yields than the Chelex® protocol.

Comparison of quantitation results for blood samples

Refer to Table 5 for observed data. Mean blood quantitation values for samples extracted using NucleoSpin® 8 Trace were comparable to Chelex® results. Yields were variable but comparable to Chelex®.

Comparison of DNA profiles

NucleoSpin® 8 Trace overall yielded higher allele counts compared to Chelex resulting in 202 reportable alleles in contrast to the 89 alleles from Chelex®-extracted samples (Table 14). NucleoSpin® 8 Trace was able to yield profiles for cell samples on denim down to 1/8 dilution, but performed poorly with FTA® samples, resulting only in a partial profile (X.X+5) for the neat cell sample. NucleoSpin® 8 Trace performed better for cells on cotton swaps, and performed moderately better for cells on rayon swaps. Profiles from cells on cotton cloth samples were comparable between the two DNA extraction methods.

Table 14. Comparison of DNA profiles for cell substrate samples extracted using either Chelex NucleoSpin® 8 Trace.

CELLA

	4	•	•	*	Y	
Me	thed:	c	h	e î		

Dilution	FTA	Cotton swebs	Rayon swabs		Cetton	Dentm
	Profile -	Profile	Sample#	Profile	Profile	Profile
Nest:	X.X+18 ·	X,X+18	R14	NSD	X.X+8	NR/NSD
			R15	NEVNSD		Market Market
		914:15 7:15	R16	NSD -	Section Sections	
4	- 光學學術論學		R17	NSD	200 FEB 12	300
DI 1/4	X,X+18	XNR+3	R10	NSD	NR+1	NRVNSD
,		注意 为100	R11	NSD	Same	100
	正言語語學			NSD	40000	here we
		A STATE OF	R13	NSD	拉斯斯克拉克	State of
D3 1/8	XX+17	X,X+3	R6	NSD	XNR+3	NRVNSD .
			R7	NSD	Carrier St.	200
		经现代表现	R8	NSD		
	1000	9	R9	NSD	839K988	100 Mark 100
Dž 1/18	NSD · · ·	NSD · .	R2	NSD	NSD .	NSD · ·
	\$100 T	17(7)	R3	NSD		
	505 Sept. 100	100	R4	NSD	THE PARTY.	
	THE CONTRACT	NEW YORK	R6	NSD · ·	· 经实际公司的	

	FTA	Cotton	Rayon sw	eas A	L'Solton	Danim
	P.ro@a	Profile	Sample#	Profite "POS	見る神経を	Profile
Nezi	X,X+6 ·	X,X+18	R14	AISD3	X-X+180Es	XX418
	建设企业	1.00	R15	X,X+18	4	
		0.000	R16	X,X+16 AI@0		
	12.00		R17	AI@D13	100	205 B
Dil 1/4	NSD	XX+18	R10	X,NR+NR/NS	NSDE S	XX-18F1
	92.00	ASOBS C	R11	X.X+3	the second	MENDRALA
		250	R12	XNR+1 S		
	The state of the		R13	NR.NR +1	D: 10	50 5K2
DØ 1/8	NSD	X,X+17	R6	X,NR+NR/NS	XNR+2	X,X+13 ·
	250	《题记题录》	R7	NR/NSD 1	经 国金融	Tributa de
		100	R8	X.NR+NR/NS	1000000000	
			R9	XNR+NR/NS		14.
DR 1/18	NSO .	NSD	R2	N9D	NSDINR	NSD .
	ME SELVE		R3	NSD		500 S. Y
	No. of the last	TANK THE	R4	NSD	1000	3-11-2
	张祖明 100000	100000	R6	X.NR+NR/NS	大文学等的	S. 25 (1733) E.

For blood samples on rayon swabs, NucleoSpin® 8 Trace profiles were comparable to Chelex®, with several partial profiles being observed in the neat and 1/8 dilutions (Table



Cass | Forensic and Scientific Services Table 15. Comparison of DNA profiles for blood substrate samples extracted using either Chelex® or NucleoSpin® 8 Trace. Cotton Dilution Profile Sample# R14 R14 X,Y+18 Neat R15 X,Y+18 R15 X,Y+18 NRVNSD R17 X,Y+18 DII 1/4 Not Uploa R10 X,Y+18 X,Y+18 R12 X,Y+18 X,Y+18 R13 Dil 1/8 X,Y+18(AI@UX,Y+18 X,Y+18 NR/NSD X,Y+18 X,Y+18 Dil 1/16 X,Y+18 X,Y+18 X,Y+18 6.7 Summary Findings from the evaluation of various forensic DNA extraction kits, compared to the inhouse Chelex® protocol, is summarised in Table 16. Queensland Government

Table 16. Summary of findings from the evaluation of five forensic DNA extraction chemistries.

				A STATE OF THE STA		
	Chelex	DNA IQ™	QIAamp [®] DNA	Charge Switch *	forensic GEM™	NucleoSpin® 8 Trace
Processing time for 12 samples	2hr	3hr	5hr			5hr
Nashing steps included to remove inhibitors	No.	Yes	Yes		No	Yes .
rinal extract volume (μL)	~150	100	45		100 for FTA, 200 for other samples	100
6 zero quantitation values for cells	43.750	9.375	37(1.30	THE STATE OF THE S	9.376	24.140
zero quantitation values for blood	0.000	0.000	00.0	0,000	0.000	0.000
ell substrate displaying highest quant value for neat cell samples	FTA	Cotton swab	C tion swab	Cotton symb	Cctton swab	Cotton cloth
otal number of reportable alleles for cells (max 576)	89	282	7.6	700 E 138	209	202
otal number of reportable alleles for blood (max 288)	234	252		288	25	264
otal number of autosomal loci exhibiting allelic imbalance (max 432)	0 1	1*	3	5	1	6^
eat cell samples on denim showed inhibition (no profile)	Yes	No	Yes	Yes	No	No
eat blood samples on rayon swabs showed inhibition (no profile)	No	No-	No	No	Yes	No
menable to automation	· No	Yes	Yes	Yes	Yes	Yes
/alidated MultiPROBE II PLUS automated protocol	No	Yes	No	No# No#	No.	No .

^{*} Five occurrences of alletic imbalance were observed in Amelogenin.

Page 27 of 3

^{*}One occurrence of allelic imbalance was observed in Amelogenin.

Out of all the chemistries tested, only the Chelex® method and *forensic*GEM™ protocols do not incorporate washing steps for the removal of inhibitors and residual proteins. This is because in these protocols, the DNA is free in solution and not immobilised on to a capture device such as magnetic beads, and therefore washing of the sample cannot be performed. Washing steps result in high quality, purified DNA extracts. As such, Chelex® and *forensic*GEM™ extracts are considered to be crude DNA extracts of suboptimal quality that may not yield the best DNA profiles due to the presence of inhibitors that can affect PGR amplification of multiple STR loci. Although the dye in denim material did not appear to result in inhibition for *forensic*GEM™ samples, only 25/288 alleles (8.7%) from blood samples could be resolved by this extraction method.

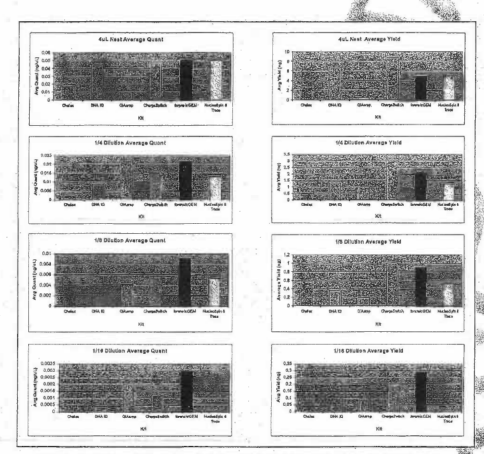


Figure 1. Average quantitation values (ng/µL) and yields (ng) for cell samples extracted using the various extraction chemistries tested, compared to Chelex.



Although all kits resulted in quantitation results for all blood samples (0% had zero results), the results for cell samples exhibited more variation. Out of the extraction chemistries that incorporate washing steps, the DNA IQ™ system exhibited the best result for zero quantitation values for both cell and blood samples at 9% and 0% respectively. Almostihalf (44%) of Chelex® cell extracts failed to yield quantitation results. The next worse quantitation results were observed for QiAamp® DNA Micro (37.5% had zero results) followed by ChargeSwitch® (31.25%) and NucleoSpin® 8 Trace (24.14%). For alk the different substrate types tested, average quantitation values were comparable for DNA QIM, ChargeSwitch®, forensicGEM™ and NucleoSpin® 8 Trace in neat, 1/4 and 1/8 dilutions (Figure 1). Compared to samples extracted using Chelex®, samples extracted using the evaluated kits displayed higher average quantitation results that were up to 7.7 times higher than Chelex® results. Chelex® and NucleoSpin® 8 Trace were the only two kits that did not result in quantitation values for the 1/16 dilutions. The average yields varied widely due to different elution volumes for the various kits. For neat samples, DNAIQ™ ChargeSwitch®, forensicGEM™ and NucleoSpin® 8 Trace resulted proomparable yields for neat samples, which were on average double the yield generated by Chelex® (Figure 1). In all experiments, forensicGEM™ resulted in the highest quantitation values, but as discussed in the previous paragraph, this kit produced the least number of reportable alleles for blood samples. It was preferred to have a high quantitation result, coupled with a high yield and high final volume as it allows multiple tests to be performed.

The relationship between quantitation result and the number of resolved reportable alleles is close to proportional. A list of the evaluated chemistries, ranked according to the lowest to highest percentage of zero quantitation results, and also the most to the least number of resolved alleles, is outlined in Table 17.

Table 17. A ranking of the evaluated kits based on quantitation and DNA profile results

I GIDIC II	. retaining of the oralidated file	backs ou deautiful out of the p	ti t bi oni o ilou o o iloi
Rank	% zero quantitation values	Total alleles for cells	Total alleles for blood
1	DNA IQ™ & forensicGEM™	DNA IQ™	Charge Switch 8
2	NucleoSpin® 8 Trace	forensicGEM TM	QIAamb DNA Micro
3	Charge Switch [®]	NucleoSpin® 8 Trace	Nucleo Spin 8 Trace
4	QIAamp® DNA Micro	ChargeSwitch [®]	DNAIGH
5	Chelex®	Chelex®	Chelex
6		OlAamp® DNA Micro	forensicGFN/TM

The DNA IQ™ system was ranked the highest for most categories and performed the best for both cell and blood samples (see also Figures 2 and 3). For blood samples on rayon swabs, DNA IQ™ received a lower ranking due to 2 outlier results for neat dilutions as discussed above, but overall was considered to produce the best result for all dilutions. In contrast, Chelex® had the lowest rating as it was found to result in the least number of reportable alleles for both cell and blood samples. *forensic*GEM™ also outperformed the other kits for cell samples but performed very poorly for neat blood samples, indicating an inhibitory effect due to dissolved heme, although PCR amplification performance was improved in extracts of diluted blood samples (Figure 3). In contrast, QIAamp® DNA Microworked well for blood samples, but performed the worst for cell samples. ChargeSwitch®, the alternative magnetic bead system to DNA IQ™, also performed better for blood samples than cell samples. The NucleoSpin® 8 Trace system, another membrane-based technology, performed moderately well and was ranked 3rd for the total number of alleles resolved for both cell and blood samples. Our results did not clearly indicate as to which technology, i.e. magnetic bead or silica membrane, was overall a better DNA extraction technology for forensic samples. However, DNA IQ™ worked the best in our hands as a complete "out-of-the-box" solution for extracting both cell and blood samples on various types of substrates.



Page 29 of 34

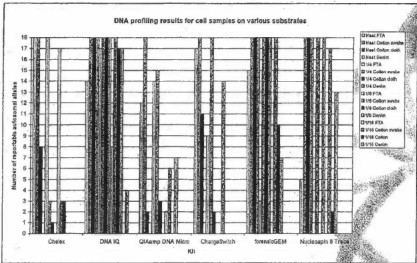


Figure 2. Total number of reportable alleles generated for cell samples on various substrates that were extracted using the various extraction chemistries tested, compared to Chelex. The kit significant is given by the most number of full bars (i.e. most full profiles) was found to be DNA QTM indicating the superior performance of this kit over the other kits tested. The current in-house Chelex method did not perform as well as several of the tested kits.

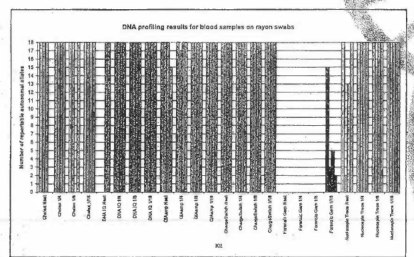


Figure 3. Total number of reportable alleles generated for blood samples on rayon swabs that were extracted using the various extraction chemistries tested, compared to Chelex. All kits were able to resolve profiles from most dilutions, except *forensic*GEM™ which could only resolve alleles from the 1/16 dilution, indicating an inhibitory effect of heme on the *forensic*GEM™ system.



age 30 of 34

Out of a total of 432 loci amplified in the assessment of each kit, only one occurrence of allelic imbalance (AI; where peak height ratio is <70%) was detected in each of the Chelex®, DNA IQ™ and forensicGEM™ kits (Table 16). QIAamp® DNA Micro and ChargeSwitch® each showed 3 and 5 occurrences of AI respectively, and NucleoSpin® Trace showed the most AI at 6 occurrences observed (Table 16).

Out of 17 occurrences of AI in all samples tested, 9 AI events were observed in cell; samples and 8 events were observed in blood samples (Table 18). These results do not suggest any increased likelihood in observing AI in either cell or blood samples. Out of the 9 AmpF(STR® Profiler Plus loci interrogated, AI was only encountered in 6 loci; D3S1358, FGA, D13S317, D8S1179, D18S51, and D7S820 (Table 18). Most of the AI (35.29%) occurred in the D13S317 locus, and the least (5.88%) occurred in both D8S1179 and D7S820. The %AI observed was within the range of 52.30% at D13S317 to 69.96% at D3S1358 (data not shown). Most of the AI (58.82%) was ≥60%, and ±1.18% of AI was ≥65%. Out of the 7 occurrences of AI that were ≤60%, 4 events (57%) were observed in cell samples extracted using NucleoSpIn® 8 Trace. Six additional occurrences of AI, were observed in Amelogenin, with all AI events ≥60% (data not shown). The AI data from this evaluation will contribute to further studies on a revised in-house AI threshold.

Table 18. Frequency of all autosomal allellc imbalance observed in the evaluation

Kit	Number of	autosomal Al		Profiler Plus loci exhibiting allelic imbalance						
	Cell	Blood	D3S1358	FGA	D13S317	D8S1179%	D18S51 D7S820			
Chalex	0	1	1			1/2/8	A CARLON			
DNA IQ	1				1					
QiAamp DNA Micro	1	2				1	2.56			
ChargeSwitch	2	3	1	1	2		1			
forensicGEM		1			1		203			
NucleoSpin 8 Trace	5	1	2	1	2	197	1,3,2,30			
Total	9	8	4	2	6	1	1 3			
		17	23.53%	11.76%	35.29%	5.88%	17.65% 5.88%			

Neat cell or blood samples that were extracted using the various kits displayed varying inhibition results for denim dye and heme (Table 16). In several cases, if a kit did not show inhibition for denim dye, it would show inhibition for heme, or vice versa. Only the DNA IQ™ and NucleoSpin® 8 Trace systems did not indicate inhibition for either inhibitor. Ther did not appear to be a link between the presence or absence of inhibition and the observation of allelic imbalance, although DNA IQ™ and NucleoSpin® 8 Trace generated the most number of total reportable alleles (534 and 466 alleles respectively). These resusuggest that the ability to remove inhibitors (such as encountered in the DNA IQ™ and NucleoSpin® 8 Trace protocols) can result in an increase in the number of resolvable alleles, therefore successfully obtaining more DNA profile results more often.

Cotton substrates (e.g. cotton swabs and cotton cloth) make up a large percentage of samples processed in DNA Analysis FSS. For example, cotton swabs make up around 45% of the total number of sample types analysed for DNA analysis (Figure 4). It was therefore considered important that the DNA extraction kits evaluated could process samples and stains on cotton matrices. It was found that the neat cell samples that displayed the highest quantitation values across all extraction kits originated from cotton swab substrates, except for Chelex® results where the best result came from FTA (Table 16).



2age 31 of 34

All of the forensic DNA extraction kits evaluated are amenable to automation, and automated protocols already exist for several kits. However, only the DNA IQ™ kit has been validated for use on the MultiPROBE® II PLUS HT EX platform and a validated protocol was developed by PerkinElmer (PerkinElmer, 2004).

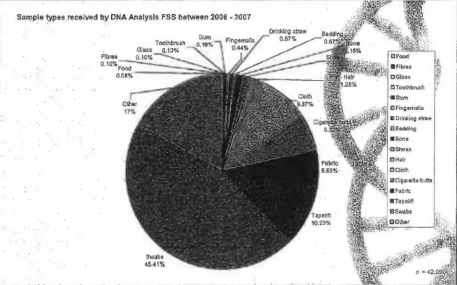


Figure 4. Pie chart of various sample types received by DNA Analysis FSS between 2006 and 2007 Around 45% of samples received for DNA analysis are swab substrates. Data was obtained from AUSLAB on 14 November 2007.

Some of the concerns raised regarding some of the kits tested include:

- QIAamp® DNA Micro involved multiple tube transfers that increased the risk of
- cross-contamination and also increased processing time to 5 hours for 12 samples. An increased risk of contamination was also prevalent in the NucleoSpin® 8 Traces method when coupled with the PVC vacuum manifold, because of the need to fit multiple adapters to ensure seals are maintained for a proper vacuum environment If the plates and adapters were not assembled correctly, the vacuum environment would fail and possibly cause cross-contamination and, more alarmingly, loss of sample. Furthermore, even when assembled correctly, biohazardous contaminants (e.g. blood) are drawn down the manifold through the vacuum tubing and into the collection containers. Decontamination of the tubing and containers raises serious health and safety concerns.
- The forensicGEM™ system was the quickest protocol to perform and yielded crude DNA extracts that produced high allele counts for cell samples. However, the system could not deal with blood samples (and heme inhibition) effectively, therefore causing very low allele counts for blood samples.
- ChargeSwitch® was the alternative magnetic bead system to DNA IQ™. However, ChargeSwitch® did not produce results that were comparable or better than DNA.



Page 32 of 34

IQ™. For example, more ChargeSwitch® samples did not yield quantitation results compared to DNA IQ™ and resulted in a lower total allele count. ChargeSwitch® also did not appear to be able to effectively deal with Inhibition from the dye in denim material.

Overall, data from the evaluation suggested that DNA IQ™ outperforms all of the forensic DNA extraction kits tested, in addition to the in-house Chelex® protocol. In summary, DNA IQ™:

- Is quick to perform the amount of time taken to complete the DNA extraction protocol is comparable to the in-house Chelex® method;
- Includes washing steps to remove inhibitors washing of the immobilised DNA enables purified DNA template to be eluted;
- Produced DNA quantitation values for most (>90%) samples the percentage of samples that did not yield a quantitation result was one of the lowest for DNA Q.T.
- Generated the highest number of total reportable alleles samples extracted using DNA IQ™ produced 65% more resolved alleles compared to cheex®;
- Exhibited minimal allelic imbalance the occurrence of Al in DNA IQ ** samples was comparable to Chelex*, although increased Al in Amelogenia was observed;
- · Was not inhibited by heme in blood samples;
- · Was not inhibited by the dye In denim material;
- Has been validated for use on the MultiPROBE® II PLUS HT EX platform.

7. Recommendations

Based on the results from evaluating various commercial DNA extraction kits that were designed specifically for forensic use, and comparing results from each kittotine current inhouse Chelex[®] protocol, we have found DNA IQ™ to be the most suitable kit for extracting cell and blood samples that are analysed in DNA Analysis FSS. We therefore recommend that further studies be performed on the DNA IQ™ system in order to:

- Validate a manual DNA IQ™ protocol for extracting various DNA Adaiysis FSS substrate types;
- Verify an automated DNA IQ™ extraction program on the MultiPROBES II PLUS HT Ex platforms for automated DNA extraction of various DNA Analysis FSS substrate types.

8. References

Butler J (2005). Forensic DNA Typing. Elsevier: Burlington, MA, USA.

Gallagher B, Hlinka V, Muharam I, Iannuzzi C, Lundie G, Nurthen T, Ientile V (2007). Mos sample creation for cell and blood samples [Laboratory Report]. DNA Analysis FSS Brisbane, QLD, Australia.

Invitrogen Life Technologies (2005), ChargeSwitch® Forensic DNA Purification Kits for purification of genomic DNA from forensic samples [PN 25-0825, Version A, 3 January 2005], Invitrogen Corporation: Carlsbad, CA, USA.

Macherey-Nagel (2003). NucleoSpin® 8 Trace: Genomic DNA from Forensic Samples [Rev 0.2, November 2003]. Macherey-Nagel: Düren, Germany.

Moss D, Harbinson S-A, Saul DJ (2003). An easily automated, closed-tube forensic DNA extraction procedure using thermostable proteinase. Int J Legal Med 117:340-349.

PerkinElmer (2004). MultiPROBE II Liquid Handling Forensic Workstation Application Guide: Automated DNA IQ™ System for Mixed Casework Sample DNA Isolation [PN 8842157]. PerkinElmer Life and Analytical Sciences: Downers Grove, IL, USA.

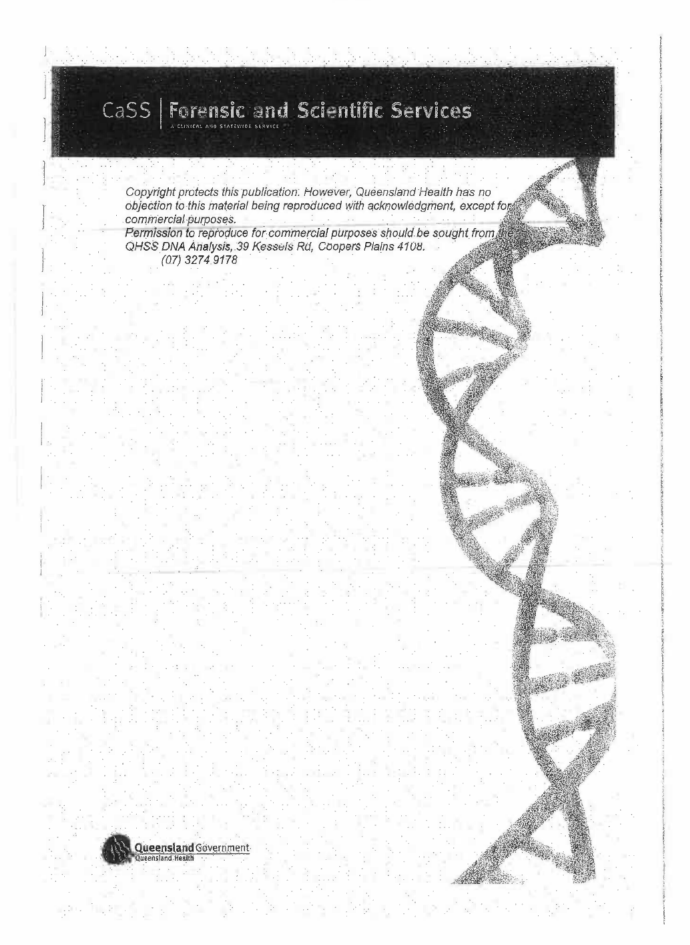
8842157]. PerkinElmer Life and Analytical Sciences: Downers Grove, IL, USA.

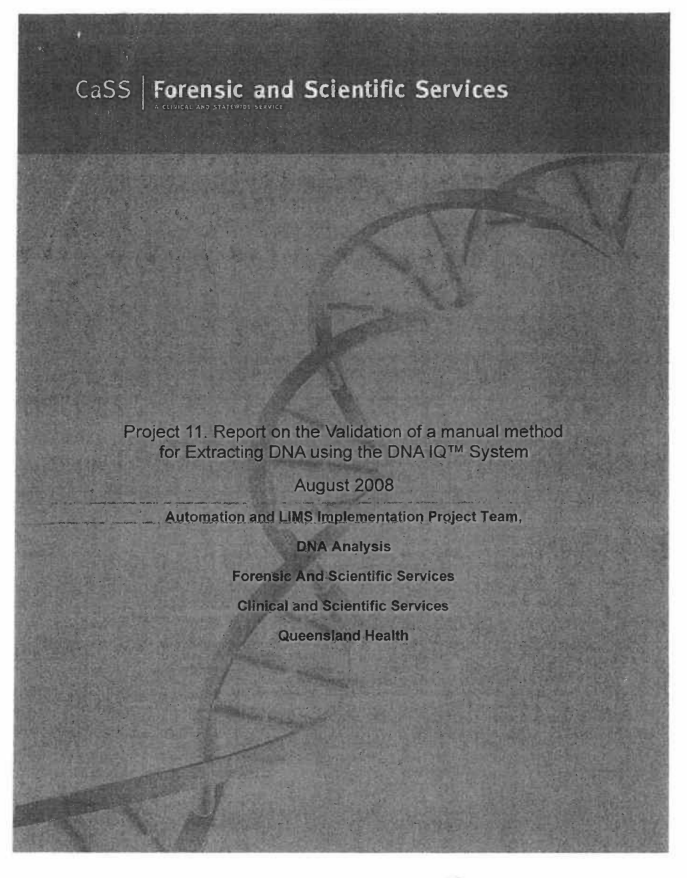
Promega Corporation (2006). DNA IQ™ System – Small Sample Casework Protocol [PN TB296, Rev. 4/06]. Promega Corporation: Madison, WI, USA.



Page 33 of 3

Cass | Forensic and Scientific Services Qlagen (2003). QlAamp® DNA Micro Handbook [Rev. 08/2003]. Qiagen GmbH: Hilden, Germany. QIS 19171 (2007). Method for Chelex® Extraction [Standard Operating Procedure]. DNA Analysis FSS: Brisbane, QLD, Australia. Scott S, Harbison S-A, Saul DJ (2003). Developmental validation of forensicGEM™: Thermophilic Proteinase for Forensic DNA Extraction [oral presentation at the International Symposium on the Forensic Sciences, Fremantle, WA, 2-7 April 2006 Institute of Environmental Research Ltd: Hamilton, New Zealand. Walsh PS, Metzger DA, Higuchi R (1991). Chelex 100 as a medium for simple extraction of DNA for PCR-based typing from forensic material. Biotechniques 10:506-13. ZyGEM (2006). DNA extraction from buccal swabs using forensicGEM. ZyGEM Corporation Ltd: Hamilton, New Zealand.





safe | sustainable | appropriate



Project 11. Report on the Validation of a Manual Method for Extracting DNA using the DNA IQ™ System

Nurthen, T., Hlinko, V., Muharam, I., Gallagher, B., Lundle, G., Iannuzzi, C., Ientile, V. Automation/LIMS Implementation Project, DNA Analysis FSS (August 2008)

Abstract

The DNA IQ™ system was found to be the most suitable kit for extracting cell and blood samples that are analysed in DNA Analysis FSS (refer to Project 9). This DNA extraction system, based on magnetic bead technology, was found to generate results that were comparable or better than the current Chelex®-100 protocol. We have validated a manual DNA IQ™ method for extracting DNA from forensic samples, and incorporated studies on sensitivity and consistency, inhibition, substrate type, substrate size, and mixture studies. This manual DNA IQ™ method is suitable for verification on the automated MultiPROBE® II PLUS HT EX extraction platforms.

2. Introduction

A previous evaluation of various DNA extraction systems that were designed specifically for forensic samples was performed in order to select a suitable extraction technology for extracting various sample types that are processed in DNA Analysis FSS. DNA IQ™ was identified as a suitable kit for extracting forensic samples, and was found to outperform both the current Chelex®-100 protocol and also all the other kits evaluated. The results of the evaluation are reported in Project 9 (Gallagher *et al.*, 2007a).

DNA purification with silica matrices, either in membrane- or bead-form, commonly uses the affinity of DNA for silica without the need for hazardous organic reagents. However, these systems tend to require extensive washing to remove the guanidium-based lysis buffer. The DNA IQ™ system uses a novel paramagnetic resin for DNA isolation (Promega Corp., 2006). The DNA IQ™ System's basic chemistry is similar to other silica-based DNA isolation technologies, except that the specific nature of the paramagnetic resin, coupled with the formulation of the lysis buffer, is unique. In the DNA IQ™ System, negativelycharged DNA molecules have a high affinity for the positively-charged paramagnetic resin under high salt conditions supplied by the lysis buffer. Once DNA is bound to the magnetic resin, and the resin is immobilised by a magnet, the sample can be washed using an alcohol/aqueous buffer mixture. The high alcohol content of the wash buffer aids to maintain the DNA-resin complex in low-salt conditions, while the aqueous component functions to wash away residual lysis buffer and any inhibitors or non-DNA contaminants such as cellular debris and protein residues. DNA is released from the resin by using a low ionic strength elution buffer, and the purified DNA can be used directly in downstream applications such as PCR.

For samples that are in excess (e.g. reference samples), DNA IQ™ resin will only isolate up to a total of approximately 100ng of DNA due to bead saturation (Huston, 2002).

3. Aim

To validate a manual method for DNA extraction of blood and cell stains on forensic samples using the DNA IQ™ system (Promega Corp., Madison, WI, USA).



Page 1 of 21

Equipment and Materials

- DNA IQ™ System (Promega Corp., Madison, WI, USA); 100 samples, Cat.# DC6701), which includes:
 - o 0.9mL Resin
 - o 40mL Lysis Buffer
 - o 30mL 2X Wash Buffer
 - 15mL Elution Buffer
- TNE buffer (10mM Tris, 100mM NaCl, 1mM EDTA, pH 8.0)
- MagneSphere® Magnetic Separation Stand, 12-position (Cat.# Z5342) (Promega Corp., Madison, WI, USA)
- DNA IQ™ Spin Baskets (Cat.# V1221) (Promega Corp., Madison, WI, USA)
- Microtube 1.5mL (Cat.# V1231) (Promega Corp., Madison, WI, USA)
- 95-100% ethanol
- Isopropyl alcohol
- 1M DTT (Sigma-Aldrich, St. Louis, MO, USA)
- Proteinase K (20mg/mL) (Sigma-Aldrich, St. Louis, MO, USA)
- 20% SDS (Biorad, Hercules, CA, USA)
- 0.9% saline solution (Baxter Healthcare, Old Toongabbie, NSW, Australia)
- ThermoMixer Comfort (Eppendorf, Hamburg, Germany)
- Vortex mixer
- Bench top centrifuge
- Cytobrush® Plus Cell Collector (Cooper Surgical, Inc., Trumbull, CT, USA)
- FTA® Classic Cards (Whatman plc, Maidstone, Kent, UK)
- Rayon (155C) and cotton (164C) plain dry swabs (Copan Italia S.p.A., Brescia, Italy)
- Vacuette® K2EDTA blood collection tubes (Greiner Bio-One GmbH, Frickenhausen, Germany)
- Sticky tape (BDF tesa tape Australia Pty Ltd)
- Tannic acid C₇₆H₅₂O₄₅ FW1701.25 (Selby's BDH, Lab Reagent >~90%)
- Urea NH₂CONH₂ FW60.06 (BDH, Molecular Biology Grade ~99.5%)
- Indigo carmine C₁₆H₈N₂Na₂O₈S₂ FW466.35 PN 131164-100G (Sigma-Aldrich, St. Louis, MO, USA)
- Humic acid sodium salt PN H167520-100G (Sigma-Aldrich, St. Louis, MO, USA)
- Used car motor oil, SW20/SAE50 (Caltex)
- Various clothing materials, including:
 - o Best & Less Pacific Cliff, White cotton shirt, XXL
 - Big W Classic Denim, Men's Blue denim jeans, 112
 - Private Encounters, off-white nylon cami, size 14
 - o Clan Laird, blue 100% wool kilt
 - Millers Essentials, blue 100% polyester camisole, size 10
 - Unknown, teal green 100% lycra swimwear
 - o Leather Belt, brown

5. Methods

5.1 Cell and blood collection

Buccal cells were collected using a modified Cytobrush® protocol (Mulot *et al.*, 2005; Satia-Abouta *et al.*, 2002). Four donors were chosen. Each donor was asked to brush the inside of one cheek for one minute. Then, with another Cytobrush®, the other cheek was also sampled. The cells collected on the brush where then resuspended in 2mL of 0.9% saline solution. Multiple collections were taken on different days.



Page 2 of 21

CaSS | Forensic and Scientific Services

Whole blood was collected from three donors by a phlebotomist as per standard collection procedures in EDTA tubes. Blood samples were refrigerated until spotting onto substrate and cell-counting step.

Table 1 lists the donor sample ID's.

Table 1. List of donor samples used for validating a manual DNA IQ™ method.

Donor ID	
Cell samples	
D1	
D2	
D3	
D4	
Blood samples	
D1	
D2	
D3	

5.2 Cell counting

Buccal cell suspensions were diluted using 0.9% saline solution to create a 1/10 dilution of the original sample prior to submitting for cell counting. All counts were performed by the Cytology Department, RBWH (QIS 15393).

Blood cell counting was performed on a 1mL aliquot of the original sample also by the Cytology Department, RBWH (QIS 15393)

5.3 Sensitivity, Reproducibility (Linearity) and Yield

Sensitivity and reproducibility of the DNA IQ^{TM} kit was assessed using dilutions of cell and blood samples.

For cell samples, dilutions were made using a sample from donor 4, diluted in 0.9% saline solution. The dilutions used were:

- Neat
- 1/10
- 1/100
- 1/1000

For blood samples, dilutions were made using a sample from donor 2, diluted in 0.9% saline solution. The dilutions used were:

- Neat
- 1/10
- 1/100
- 1,100

Mock samples were created from rayon and cotton swabs using the above dilutions. The swab heads were removed from the shaft using sterilised scalpel and tweezers. Swab heads were then cut into quarters and each quarter was then added to separate sterile 1.5mL tubes. To each quarter swab, 30µL of each neat sample or dilution was added to



Page 3 of 21

create a total of five replicates. Samples were dried using a ThermoMixer set at 56°C over 2 hours in a Class II biohazard cabinet.

5.4 Inhibition challenge

Quartered cotton swabs in sterile 1.5mL tubes were spotted with 30µL of neat cell suspension and were dried after each addition on a ThermoMixer as described previously. Neat blood samples were also created using the same method.

All the inhibitors except for the motor oil were obtained in powder form. Before making any liquid solution of the powdered inhibitors, research was conducted to determine the likely level of each inhibitor normally encountered in the environment (Hlinka *et al.*, 2007). Each solution was made at concentrations based on the information obtained (Table 2).

Table 2. Concentrations of various inhibitors used in the inhibition study.

Inhibitor	Excess/Neat Solution	Mass	Volume H₂O	Final inhibitor concentration		
Tannic acid	Excess	600mg	500µL	0.705M		
	Neat	200mg	500µL	0.235M		
Humic acid	Excess	1g	5mL	20% (w/v)		
	Neat	0.1g	5mL	2% (W/V)		
Indigo carmine	Excess	0.479	10mL	100mM		
	Neat	0.047g	10mL	10mM		
Urea	Excess	0.06g	1mL	1M		
	Neat	0.021g	1mL	0.33M		

A total of 30µL of each solution containing specified concentrations of various inhibitors was applied to the buccal cell and blood swabs prepared above. The only exception was motor oil, where only 15µL was added to the cell and blood swabs respectively. Each inhibitor sample was replicated in quadruplicate and left to dry overnight in a Class II biohazard cabinet.

To another set of prepared cell and blood swabs, an excess of each inhibitor was applied in quadruplicate for each inhibitor and allowed to dry overnight. This process was achieved by applying another solution of inhibitor exceeding the normal level (Hlinka et al., 2007).

5.5 Substrates

Swabs

Four cotton and four rayon swab quarters in sterile 1.5mL tubes were loaded with 30µL of neat cell or blood sample and were extracted once the sample had dried on the swab.

Tapelifts

Two donors were sampled using the tape most commonly used within the laboratory (BDF tesa tape). Strips of tape were firmly applied to the inside of the fore arm and lifted off. This process was then repeated until the tape was no longer adhesive. The tape was wrapped around sticky-side-in, forming a cylinder shape, and placed in a sterile 1.5mL tube. These samples were created in quadruplicate. Tape was not used as a substrate in the blood validation.

Fabric

The material types tested included:

· Denim jeans;



Page 4 of 21

CaSS | Forensic and Scientific Services

- · White 100% cotton shirt;
- Blue 100% wool kilt;
- Teal green 100% lycra swimwear;
- White 100% nylon camisole;
- · Blue 100% polyester camisole; and
- Brown 100% woven leather belt.

All material types except leather were sampled and ten 2.5cm x 2.5cm pieces were cut from each material and washed in 10% bleach following an in-house washing method to remove any contaminating DNA from outside the laboratory (Gallagher et al., 2007b). As for the leather, one strand of the leather weave was cut from the belt and washed following the same method. Once dry, the material was then cut into 0.5cm x 0.5cm pieces using sterile techniques, placed in 1.5mL tubes and 30µL of both cell sample and blood was applied to separate pieces. Each substrate sample was created in quadruplicate and dried on a ThermoMixer set at 56°C over 2 hours in a Class II biohazard cabinet.

Gum

Two types of chewing gum were chosen: (1) Wriggley's Extra White (peppermint flavour) and (2) Wriggley's Extra Green (spearmint flavour). The donor was asked to chew the gum for 30 minutes and dispose of the gum into a clip-seal plastic bag. The gum was then air dried in a Falcon tube overnight before it was frozen for roughly an hour before cutting into 3mm x 3mm x 3mm pieces and placed into sterile 1.5mL tubes. Gum substrates were not assessed for blood samples.

Cigarette butts

Two brands of cigarettes were smoked all the way through and then the butts collected. The filter paper of the butt was cut into 0.5mm² pieces and placed into sterile 1.5mL tubes. Cigarette butts were not assessed for blood samples.

FTA[®] Classic Card punches

Eight sterile 1.5mL tubes, each containing four 3.2mm FTA® Classic Card punches, were spotted with 30µL of cells or blood before being dried on a ThermoMixer. Four replicates contained sample from one donor, the other remaining four replicate tubes had a different donor sample added.

5.6 Mixture studies

Buccal cells and whole blood were obtained from a male and female donor. Dilutions were made using 0.9% saline solution to ensure that the cell concentration was equal. Dilutions were then performed on the male sample to obtain the correct ratios.

Mock samples were created using the following ratios of female to male:

- 1:1,
- 1:2,
- 1:10,
- 1:25.
- 1:50 and
- 1:100.

A total of 30µL of the female component was spotted first on to a quarter of a cotton swab in a sterile 1.5mL tube and dried on a ThermoMixer before adding another 30µL of the male component. Samples were created in quadruplicate for all ratios, for both cell and blood samples



Page 5 of 21

5.7 Substrate size

Various sizes of material were cut from a white cotton shirt:

- 0.5cm x 0.5cm,
- 1cm x 1cm,
- 2cm x 2cm.

Each piece of material was stored in individual, sterile 1.5mL tubes and 30µL of cell sample was added to the material and allowed to dry on a ThermoMixer. The same process was followed for blood samples. Five replicates were made for each sample type.

5.8 Extraction using the DNA IQ™ System (Promega Corp.)

The manual DNA IQ™ method used was based on an automated protocol developed by the Centre of Forensic Sciences (CFS) in Toronto, Ontario (PerkinElmer, 2004). A Proteinase K – SDS Extraction Buffer was made as per the recommended protocol. The 1x Extraction Buffer for one sample consisted of:

277.5µL TNE buffer 15µL Proteinase K (20mg/mL) 7.5µL 20% SDS

The TNE buffer consisted of:

1.211g Tris (10mM Tris) 2mL 0.5M EDTA (1mM EDTA) 5.844g NaCl (100mM NaCl)

The adapted manual DNA IQ™ protocol is described below:

- 1. Set one ThermoMixer at 37°C and another at 65°C.
- Ensure that appropriately sized samples are contained in a sterile 1.5mL tube. For every sample, prepare three set of labelled tubes: spin baskets (for every tube except the extraction control), 2mL SSI tubes and Nunc™ tubes.
- Prepare Extraction Buffer and add 300µL to each tube. Close the lid and vortex before incubating the tubes at 37°C on the ThermoMixer at 1000rpm for 45 minutes.
- 4. Remove the tubes from the ThermoMixer and transfer the substrate to a DNA IQ™ Spin Basket seated in a labelled 1.5mL Microtube using autoclaved twirling sticks. Then transfer the liquid to a labelled 2mL SSI sterile screw cap tube.
- Centrifuge the spin basket on a benchtop centrifuge at room temperature for 2 minutes at its maximum speed. Once completed, remove the spin basket and collect the remaining solution and pool with the original extract in the 2mL SSI sterile screw cap tube, then vortex.
- 6. Add 550 µL of Lysis Buffer to each tube.



Page 6 of 21

CaSS | Forensic and Scientific Services

- Dispense 50µL of DNA IQ™ Resin Lysis Buffer solution (7µL Resin in 43µL Lysis Buffer) to each tube. Invert the resin tube regularly to keep the beads suspended while dispensing to obtain uniform results.
- Vortex each tube for 3 seconds at high speed then place in a multitube shaker set at 1200rpm to incubate at room temperature for 5 minutes.
- 9. Vortex each tube for 2 seconds at high speed before placing the tubes in the magnetic stand. Separation will occur instantly.

Note: If resin does not form a distinct pellet on the side of the tube, or if the pellet has accidentally mixed with the solution while in the stand, vortex the tube and quickly place back in the stand.

- 10. Carefully remove and discard all of the solution without disturbing the resin pellet on the side of the tube. If some resin is drawn up in tip, gently expel resin back into tube to allow re-separation.
- 11. Remove the tube from the magnetic stand; add 125µL of prepared Lysis Buffer and vortex for 2 seconds at high speed.
- Return tube to the magnetic stand, allow for separation and then remove and discard the Lysis Buffer.
- Remove tube from the magnetic stand; add 100μL of prepared 1X Wash Buffer and vortex for 2 seconds at high speed.
- 14 Return tube to the magnetic stand, allow for separation and then remove and discard all Wash Buffer.
- 15. Repeat Steps 13 to 14 two more times for a total of three washes. Be sure that all of the solution has been removed after the last wash.
- 16. In a biohazard cabinet, place the lids of the tubes upside down on a Kimwipe, in their respective order, and the tubes into a plastic rack, and air-dry the resin for 5-15 minutes at room temperature. Do not dry for more than 20 minutes, as this may inhibit removal of DNA. Once dry, screw on the lids.
- To each samples then add 50µL of Elution Buffer very gently on the top of the magnetic pellet. Do not mix.
- 18. Close the lid and then incubate the tubes in the ThermoMixer at 65°C for 3 minutes with no shaking and another 3 minutes shaking at 1100 rpm.
- 19. Remove the tubes and vortex for 2 seconds at high speed. Immediately place the tube in the magnetic stand. Tubes must remain hot until placed in the magnetic stand or yield will decrease.
- Carefully transfer the supernatant containing the DNA to the respective labelled Nunc[™] tubes.



Page 7 of 21

CaSS | Forensic and Scientific Services

 Repeat step 17 to 20, transferring the supernatant to the appropriate Nunc[™] tube. The final volume after the second elution should be approximately 95µL.

Note: DNA can be stored at 4°C for short-term storage or at -20 or -70°C for long-term storage.

5.9 DNA quantitation

All DNA extracts were quantified using the Quantifiler™ Human DNA Quantitation kit (Applied Biosystems, Foster City, CA, USA) as per QIS 19977. Reaction setup was performed on the MultiPROBE® II PLUS HT EX (PerkinElmer) pre-PCR platform.

5.10 PCR amplification

DNA extracts were amplified using the AmpFlSTR® Profiler Plus® kit (Applied Biosystems, Foster City, CA, USA) as per QIS 19976. Reaction setup was performed on the MultiPROBE® II PLUS HT EX (PerkinElmer) pre-PCR platform.

5.11 Capillary electrophoresis and fragment analysis

PCR product was prepared for capillary electrophoresis using the manual 9+1 protocol (refer to Project 15 and QIS 19978). Capillary electrophoresis was performed on an ABI Prism® 3100 Genetic Analyzer (Applied Biosystems, Foster City, CA, USA) under the following conditions: 3kV injection voltage, 10 sec injection time, 15kV run voltage, 100µA run current, and 45min run time. Data Collection Software version 1.1 was used to collect raw data from the ABI Prism® 3100 Genetic Analyzer. Fragment size analysis was performed using GeneScan 3.7.1. Allele designation was performed using Genotyper 3.7, with thresholds for heterozygous and homozygous peaks at 150 and 300 RFU respectively. The allelic imbalance threshold is 70%.



Page 8 of 21

6. Results and Discussion

6.1 Donor sample cell counts

Aliquots of buccal cell samples were counted at Cytology Department (RBWH) to determine the concentration of viable cells, in order to better estimate the number of cells at any particular dilution. A white cell count was not performed on all the blood samples, and therefore an estimate on the number of nucleated cells could not be determined.

6.2 Sensitivity, consistency and yield

To ensure the reliability and integrity of results for samples containing small amounts of DNA, a sensitivity study was conducted to determine the lowest concentration of DNA that provides reliable results. A consistency study was combined into the sensitivity experiment to determine the maximum acceptable difference between the results obtained. All samples were extracted in identical conditions by the same operator at the same time to minimise variability.

The cell sample used for the experiments was from donor sample 4A, which was counted to be around 3,680 nucleated cells (x 10^6 /L). The blood sample used was from donor 6A, which was counted to be around 2,540 nucleated cells (x 10^6 /L). The estimated amount of DNA present in each dilution is outlined in Table 3.

Table 3. Amount of DNA in each dilution, as calculated from the cell count.

Sample type	Dilution factor	Number of cells (/µL)	gDNA (ng/μL)	Theoretical total DNA on swab (ng)
	Neat	3680	23.552	706.56000
Calla	1/10	368	2.3552	70.65600
Cells	1/100	36.8	0.23552	7 06560
	1/1000	3.68	0.023552	0.07656
	Neat	2540	16.256	487.68000
Disad	1/10	254	1.6256	48.76800
Blood	1/100	25.4	0.16256	4.87680
	1/1000	2.54	0.016256	0.48768

The DNA yields resulted from extracting the above cell dilutions using the DNA IQ™ System is outlined in Table 4. Blood samples produced higher yields compared to cell samples. On average, blood samples on cotton swabs generated the highest yields. Cell samples on rayon and cotton swabs generated similar yields. All blood dilutions down to 1/1000 produced quantitation results, but cell samples only produced reliable quantitation results down to 1/100 dilution, possibly due to the effects of cell clumping.



Page 9 of 21

Table 4 DNA quantitation data for diluted cell and blood samples on rayon and cotton substrates.

Sample type	Dilution factor	Theoretical Input DNA (ng)	Rayon swab yield (ng)	Alleles	Cotton swab yield (ng)	Alleles	Rayon average yield (ng)	Rayon Std Dev	Recovery Rayon (%)	Cotton average yield (ng)	Cotton Std Dev	Recovery Cotton (%)
			110.0000	18	117.0000	18					S.Fr. 3-3	
			130 0000	18	124.0000	18					CINCOLOR -	
	Neat	706,56000	160.0000	18	46.8000	18	134.5400	41.30	19.04	95.2800	32.69	13.48
			83 7000	7	76.6000	18						
			189.0000	17	112 0000	18			5400PC-1	aff.		
			10.1000	18	12.8000	18			43400	65%		
			12.7000	18	6.3100	18						
	1/10	70 65600	9.5500	18	11,5000	18	10.4520	1.44	14.79	10.4820	2.52	14.84
			9.0100	18	10.1000	18			307 "72			
Cells			10,9000	18	11,7000	18			490"			
Cells			0.6350	0	0 0000	0			USTRA			
			0.4930	0	0.0000	0			STOCKED TO STOCK	788		
	1/100	7 06560	1.4000	5	0 2770	0	0.9254	0.64	13.10	0.1270	0.18	1.80
			1.7900	14	0 3580	0						
			0.3090	0	0.0000	0						
			0.0000	0	0.3630	0			LC ACTION			
			0.0000	0	0 0000	0			SSSS (000)			
	1/1000	0.7656	0.0000	0	0.0000	0	0.0166	0.04	2.17	0.0726	0.16	9.48
			0.0831	0	0.0000	0			25000** A606**			
			0.0000	0	0 0000	0			100, 407			
			216.0000	18	718.0000	18			TOTAL 4507			
			447.0000	18	297.0000	18			1000000 -			
	Neat	487.68000	215.0000	18	595.0000	18	317,0000	102.36	65.00	447,0000	196.46	91 66
			383.0000	7	326_0000	18			HMESA.			
			324.0000	18	299 0000	18			0075000			
			113.0000	18	126,0000	18						
			107.0000	18	91,9000	18			FG 75			
	1/10	48.76800	145 0000	18	75,4000	18	124,7800	28.10	255.86	97.6600	21.66	200.25
		1011 0000	95,9000	18	81.0000	18			IOD Have the A			
			163.0000	18	114.0000	18			Spinister of the last of the l	200		
Blood			14.3000	18	15.9000	18			7770			
			12.5000	13	12 1000	18				~ (SECTION)		
	1/100	4.87680	13.2000	18	20.8000	18	12.4800	1.62	255,91	16,7600	4.69	343.67
		1101000	9,9000	18	22,4000	18						
			12.5000	18	12.6000	18						
			0.7300	18	2,3700	18			750			
			0.6990	18	3.1300	18			7000	ART 1557		
	1/1000	0 48768	1.1800	18	3.6300	18	0 8894	0.20	182.37	3.0200	0.85	619.26
			0.8670	18	1.9700	18			Villa	VOT AND		
			0.9710	18	4.0000	18			701	200. 2000		

Page 10 of 21

The average yield observed within cell and blood samples on either rayon or cotton swabs were comparable (Figure 1). Some inconsistencies were present in cell samples at the lower dilutions of 1/100 and 1/1000 due to unreliable quantitation data at these low dilutions. Blood samples were found to generate higher average yields than cell samples and gave unexpectedly higher recovery values, despite the fact that the input DNA amount was 2-fold higher for cells compared to blood samples (Table 4). This discrepancy may have arisen from inconsistencies in cell suspension uniformity during dilutions of the original cell or blood sample, resulting in inaccurate estimates for average cell concentrations.

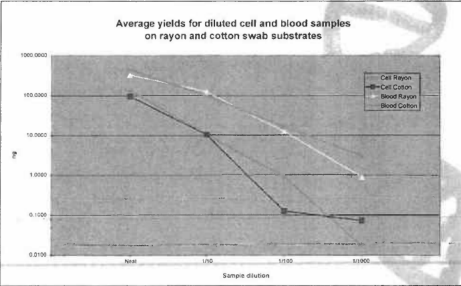


Figure 1. Average yields as observed in the sensitivity study. The yields for cell and blood samples, on two different swab types, were comparable as indicated by overlapping lines on the graph.

The dilution factor was, however, accurately reflected in the average yield for the various dilutions as displayed in Table 4 and Figure 2. An exception to this was the average yields for the neat dilutions (Figure 2). DNA IQ™ isolates a maximum of 100ng DNA as the resin is present in excess, and the system becomes more efficient with samples containing less than 50ng of DNA. Because the amount of DNA was in excess in neat samples, the observed yields varied from sample-to-sample. According to the manufacturer, the DNA IQ™ Database Protocol should be used for samples containing more than 100ng DNA to result in more consistent concentrations between the samples (Huston, 2002).

All five replicates for each neat dilution displayed the highest yields for each dilution series, as expected (Figure 2). For blood samples on rayon and cotton swabs, yields were still around 1ng for samples at the 1/1000 dilution (Figure 3).



Page 11 of 21

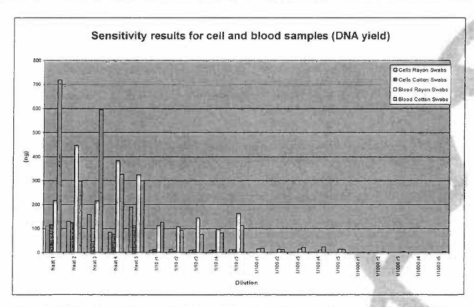


Figure 2. DNA yields (ng) observed for the sensitivity study. As expected, neat samples provided the highest yields. Yields were obtained down to 1/1000 for blood samples and 1/100 for cell samples.

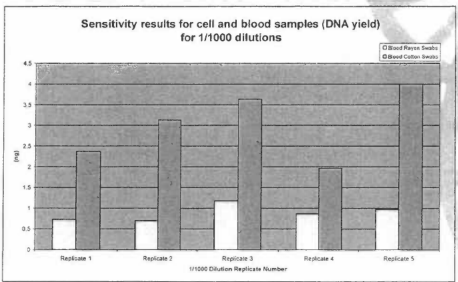


Figure 3. DNA yields (ng) observed for the sensitivity study, at the 1/1000 dilution.



Page 12 of 21

When amplified using a 9-locus STR system, all neat samples produced the expected full DNA profile (18/18 alleles), although one outlier was encountered for a cell rayon sample which produced a 7/18 partial profile (Table 4). For cell samples, full profiles could be obtained for samples that were diluted down to 1/10, with partial profiles generated from samples diluted to 1/100. For blood samples, full profiles were generally obtained from all dilutions down to 1/1000. Although two partial profiles were encountered in blood samples on rayon swabs, all blood cotton swabs produced full profiles at all dilutions.

The apparent discrepancy between the results for cell and blood samples can be attributed to inaccurate cell counts or non-uniform sample suspensions when creating the dilutions, as caused by cell clumping or cellular breakdown and precipitation.

For five replicates of each dilution, consistency was observed to vary depending on the dilution (Figure 4). Consistency, as an indication of reproducibility, was calculated as the percentage of the yield standard deviation over five replicates divided by the mean yield of all five replicates (%[\$SD_{rield}\$ / mean_yield]). A value closer to 0% indicates minimal sample-to-sample variation and therefore the results are highly consistent. The mean combined reproducibility for all neat, 1/10, 1/100 and 1/1000 dilutions were 35.31%, 20.63%, 62.14% and 124.32% respectively (Figure 4), indicating that there was high reproducibility between the neat and 1/10 dilutions across the four sample types, and reduced reproducibility at the lower 1/100 and 1/1000 dilutions. Overall, the blood samples on rayon and cotton both exhibited high reproducibility across all dilutions at an average of 30.54% and 22.45% respectively (Figure 5). The cell rayon and cotton samples were more variable across all dilutions, producing lower reproducibility at an average of 84.23% and 105.19% respectively (Figure 5). The poor performance of the cell samples can be attributed to inconsistencies in quantitation data observed at the lower 1/100 and 1/1000 dilutions.

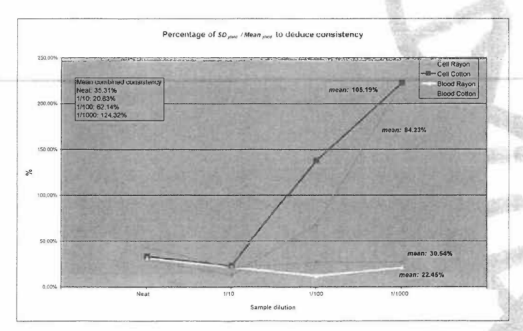


Figure 4. Reproducibility between replicates for cell and blood samples diluted down to 1/1000.



Page 13 of 21

6.3 Inhibition

Forensic samples that are commonly submitted for DNA analysis often contain inhibitors. These inhibitors may inhibit or significantly reduce the efficiency of a DNA extraction system, either by interfering with cell lysis or interfering by nucleic acid degradation or capture, therefore manifesting as extraction inhibitors (Butler, 2005). Inhibitors can also coextract with the DNA and inhibit downstream PCR amplification processes, therefore acting as PCR inhibitors (Butler, 2005). For example, inhibitors such as hemoglobin and indigo dye likely bind in the active site of the *Taq* DNA polymerase and prevent its proper functioning during PCR amplification.

For the inhibition study, five substances were chosen for their known ability to inhibit PCR and their likelihood of appearing in routine casework samples:

- Indigo carmine: a component of the blue-dye encountered in denim jeans (Shutler, et al., 1999).
- Tannic acid: a chemical used in the leather tanning process.
- Urea: a component of urine (Mahony et al., 1998).
- Humic acid: a component found in soil and soil products (Tsai and Olson, 1992).
- Motor oil: contains various hydrocarbons and ethanolic compounds that can inhibit PCR.

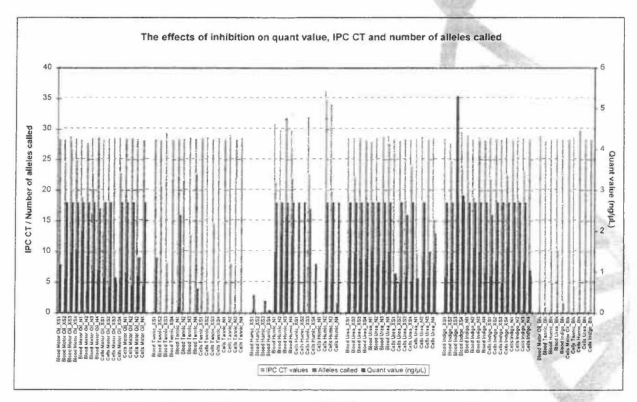


Figure 5. Effects of various inhibitors on quant value, IPC CT and number of resolved alleles.



Page 14 of 21

The observed effects of these inhibitors at neat and excess concentrations on the ability to extract, quantify and amplify various DNA samples are graphed in Figure 5. Samples were quantified using the Quantifiler™ Human DNA Quantitation Kit (Applied Biosystems) as this kit includes a built-in inhibition detector. Reaction efficiency and the presence of inhibitors can be assessed based on the performance of the internal positive control (IPC), which is known to be detected in this laboratory at around 28 cycles.

The observations that were made include:

- Samples that were spiked with motor oil, urea and indigo carmine dye did not show inhibition as determined by the IPC, and resulted in quantifiable DNA templates after extraction using DNA IQ™. The average DNA concentration observed for all samples was around 1ng/µL. The majority of samples yielded full DNA profiles, with the exception of several cell samples that were treated with urea (both at excess and neat concentrations).
- Blood and cell samples that were spiked with tannic acid did not show inhibition in Quantifiler™, as the IPC performed as expected. However, almost no amplifiable template DNA could be quantified and the majority of samples did not produce DNA profiles. This suggests that the original template DNA was degraded by application of tannic acid to the sample. It should be mentioned at this point that the tannic acid used was in the form of a yellow-brown paste substance that was applied directly to the sample swabs. The tannic acid paste, even at the neat concentration, may have been strong enough to severely fragment DNA to result in non-amplifiable templates. It was observed that three blood samples (1 with tannic acid in excess and 2 with tannic acid at neat concentration) yielded partial profiles (between 4-16 reportable alleles), and none of the cell samples produced reportable alleles. This may be caused by: (1) the concentration of viable cells in the buccal cell samples was lower than blood samples; (2) the drying of the blood stain on the substrate may have created a better barrier to protect the blood components from the degradative effects of the tannic acid.
- Blood and cell samples that were treated with humic acid in excess appeared to retain inhibition after extraction using DNA IQTM. However, at neat concentration, the effect of the humlc acid Inhibitor was overcome and amplifiable DNA template was purified as demonstrated by high DNA concentration yields. Residual inhibition was still present at neat concentration, as evidenced by higher CT values for the IPC (closer to 30), but full profiles were still produced. For some cell samples with humic acid in excess, the QuantifilerTM data suggested full inhibition (undetermined IPC CT and quantitation results), but two samples resulted in full DNA profiles.
- All reagent blanks were undetermined, indicating the absence of contamination in the results.

The results show that the DNA IQ™ system could be used to extract blood or cell samples that were spiked with motor oil, urea and indigo carmine at both excess and neat concentrations. Blood samples that contained humic acid in excess did not yield amplifiable template DNA, but 2 out of 4 cell samples with humic acid in excess appeared to produce full profiles. Samples that were exposed to tannic acid, at both neat and excess concentrations, resulted in non-amplifiable DNA, but the inhibitor was effectively washed out of the extract by DNA IQ™ as evidenced by the amplification of the IPC at the expected CT. Based on these results, we conclude that the DNA IQ™ system effectively removes inhibitors that are present in the original sample, resulting in a DNA extract that is of sufficient quality and is suitable for PCR amplification.



Page 15 of 21

6.4 Substrates

The substrate types examined included: swabs (cotton and rayon), tapelifts, fabric (denim, cotton, wool, lycra, nylon, polyester, leather), gum, cigarette butts, and FTA[®] paper. Celt and blood materials were spotted on to the substrates and extracted using DNA IQ™. The results for the two different sample types are presented in Figures 6 and 7 below.

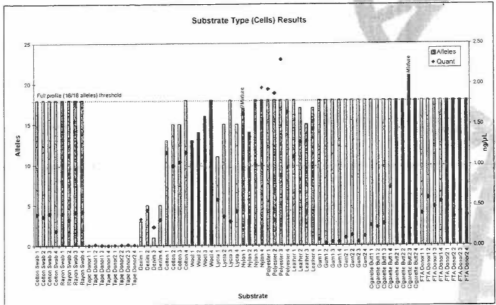


Figure 6. Number of reportable alleles and quantitation results for different substrate types containing



Page 16 of 21

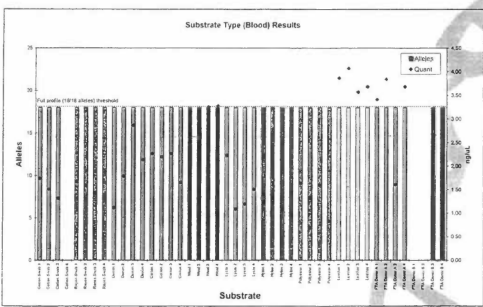


Figure 7. Number of reportable alleles and quantitation results for different substrate types containing blood material.

For cell samples:

- Full DNA profiles (18/18 alleles) were obtained from samples on cotton and rayon swabs, gum, cigarette butts and FTA[®] paper.
- The quantitation results for most of these samples were less than 0.5ng/µL. For gum samples, the average quantitation result was 0.072ng/µL, and therefore a PCR amplification at maximum volume (20µL) resulted in a total input DNA amount of 1.44ng which is sufficient to result in a full DNA profile.
- Tapelift samples gave an average quantitation result of 0.006ng/µL (just 0.002ng/µL higher than the observed background), and yielded no reportable alleles at all.
- The performance of clothing substrates was variable.
 - Cells on denim yielded quantitation results less than 0.5ng/µL but only partial profiles (maximum 5 reportable alleles), although Quantifiler™ results did not indicate any inhibition of the IPC. The poor performance of these samples may have been a result of sample preparation due to cell clumping.
 - Cells on cotton, wool and nylon resulted in higher quantitation values than lycra, but all substrates generated a similar number of reportable alleles (mean = 14 alleles). Only 25% of samples generated full DNA profiles.
 - Three out of four samples on polyester produced high quantitation results (~2ng/µL) but all samples resulted in a full profile.
 - Cells on leather displayed an average quantitation result of 1.3ng/μL and generated more than 15 reportable alleles.



Page 17 of 21

CaSS | Forensic and Scientific Services

For blood samples:

- All substrate types generated full DNA profiles.
- On average, the DNA quantitation results for all blood samples was greater than
 those resulted from cell samples. This is as per expected and was observed
 previously (see Project 9 report), because the concentration of nucleated cells in
 the blood samples were hypothesised to be higher than the concentration of buccal
 cell samples.
- Because of processing error, data was not available for the following samples.
 Cotton Swab 4, FTA Donor B 1 and FTA Donor B 2.

The results above are initial amplification results that do not take into account any reworking options.

We found that samples on tapelift substrates performed the worst; however this was probably due to the sampling method devised for this experiment, which did not adequately sample a sufficient number of cells.

6.5 Mixture studies

A mixture study was performed as part of the validation, however the results are not presented in this document because the mixture ratio was found to be inaccurate because cell counts were not performed on the saliva samples. Therefore, little information could be deduced from these results.

6.6 Substrate size

Blood on cotton swabs produced full DNA profiles for all sample sizes, ranging from $0.5\,\mathrm{x}$ $0.5\mathrm{cm}$ to $2.0\,\mathrm{x}$ $2.0\mathrm{cm}$ (Figure 8). Cells on cotton swabs did not perform as well (Figure 8), possibly due to the nature of the cells and difficulties in obtaining full DNA profiles from cell samples as observed in previous experiments.

Although the same starting amount of sample was used, it was observed that the 0.5 x 0.5cm samples generated higher quantitation results (therefore, also higher yields) compared to the 2.0 x 2.0cm samples (Figure 8). It appears that extraction efficiency decreases as the substrate surface area increases. This may be due to insufficient mixing and distribution of the lysis buffer over a larger substrate surface area, causing insufficient lysis of cellular material. This observation is in line with other reports that the DNA IQ $^{\text{TM}}$ system works more efficiently with smaller samples (Promega, 2006). The resulting IPC CT fell within the narrow range of 27.91 – 28.43 (mean = 28.10), indicating that both small and larger samples resulted in DNA extracts of similar quality, but the overall yield was lower for larger substrates (Figure 8 & 9).



Page 18 of 21

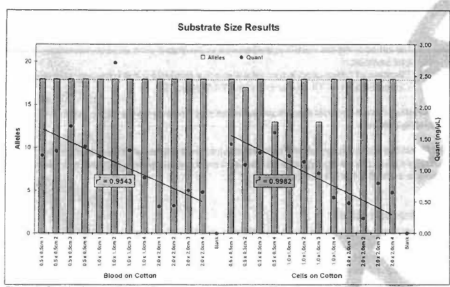


Figure 8 Results for blood and cell samples on cotton substrates of various sizes. All blood samples generated full profiles, but cell samples were more variable. The quantitation results for 0.5×0.5 cm samples were higher than those for 2.0×2.0 cm samples (blood $r^2 = 0.9543^*$; cell $r^2 = 0.9982$; *Note: an outlier was removed from the calculation).

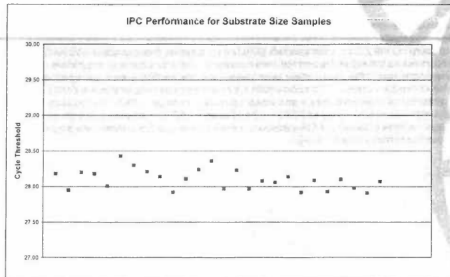


Figure 9. Various sample sizes resulted in similar CT values for the IPC, indicating that IPC performance is not affected by sample size, and that one sample size does not display a level of inhibition that is different to another sample size.



Page 19 of 21

CaSS | Forensic and Scientific Services

7. Summary and Recommendations

Based on the findings of this validation report, we recommend:

- To enable processing of cell and blood samples using the validated manual DNA IQTM protocol, except for samples on tapelift substrates.
- To design and verify an automated protocol of the validated DNA IQ[™] method for use on the MultiPROBE[®] II PLUS HT EX platforms, for processing blood and cell samples.

8. Acknowledgements

We wish to thank the Cytology Department at the Royal Brisbane and Women's Hospital for assistance with the cell-counting protocols.

9. References

- Butler JM (2005). Forensic DNA Typing: Biology, Technology, and Genetics of STR Markers, 2nd Edition. Elsevier: Burlington, MA, USA.
- Gallagher B, Hlinka V, Iannuzzi C, Lundie G, Muharam I, Nurthen T, Ientile V (2007a).

 Project 9: Report on the Evaluation of Commercial DNA Extraction Chemistries [Laboratory Report]. DNA Analysis, FSS: Brisbane, QLD, Australia.
- Gallagher B, Hlinka V, Muharam I, Iannuzzi C, Lundie G, Nurthen T, Ientile V (2007b). Mock sample creation for cell and blood samples [Laboratory Report]. DNA Analysis FSS: Brisbane, QLD, Australia.
- Hlinka V, Gallagher B, Iannuzzi C, Lundie G, Muharam I, Nurthen T, Ientile V (2007). Evaluation of the effect of inhibitors as assessed by a silica-based DNA extraction method and real-time PCR [in preparation].
- Huston K (2002). DNA IQ™ System "Frequently Asked Questions". *Profiles in DNA* 5(1):11-12.
- Mahony J, Chong S, Jang D, Luinstra K, Faught M, Dalby D, Sellors J and Chernesky M (1998). Urine specimens from pregnant and nonpregnant women inhibitory to amplification of *Chlamydia trachomatis* nucleic acid by PCR, ligase chain reaction, and transcription-mediated amplification: identification of urinary substances associated with inhibition and removal of inhibitory activity. *Journal of Clinical Microbiology* 36:3122-3126.
- Muharam I, McNevin A, Iannuzzi C, Nurthen T, Ientile V (2007). Project 15: Report on the verification of automated capillary electrophoresis setup using the MultiPROBE® II PLUS HT EX platform [Laboratory Report]. DNA Analysis, FSS: Brisbane, QLD, Australia.
- Mulot C, Stücker I, Clavel J, Beaune P, Loriot M-A (2005). Collection of human genomic DNA from buccal cells for genetic studies: comparison between Cytobrush, mouthwash and treated card. *Journal of Biomedicine and Biotechnology* 2005(3):291-6.
- PerkinElmer (2004). MultiPROBE II Liquid Handling Forensic Workstation Application Guide: Automated DNA IQ™ System for Mixed Casework Sample DNA Isolation [PN 8842157]. PerkinElmer Life and Analytical Sciences: Downers Grove, IL, USA.
- Promega Corporation (2006). DNA IQ™ System Small Sample Casework Protocol [PN TB296, Rev. 4/06]. Promega Corporation: Madison, WI.
- QIS 15396 R4 (2007). Method for bronchoalveolar lavage (BAL) cell count and differential cell count. [Standard Operating Procedure]. Anatomical Pathology, RBWH: Brisbane, Australia



Page 20 of 21

CaSS | Forensic and Scientific Services

- QIS 19976 R1 (2007). Automated amplification of extracted DNA using the AmpFtSTR® Profiler Plus® or AmpFtSTR® COfiler® kit [Standard Operating Procedure]. DNA Analysis FSS: Coopers Plains, Brisbane, Australia.
- QIS 19977 R1 (2007). Automated quantification of extracted DNA using the Quantifiler™ Human DNA Quantification Kit [Standard Operating Procedure]. DNA Analysis, FSS: Coopers Plains, Brisbane, Australia.
- QIS 19978 R0 (2008). Capillary electrophoresis setup [Standard Operating Procedure]. DNA Analysis, FSS: Coopers Plains, Brisbane, Australia.
- Satia-Abouta, King IB, Abouta JS, Thornquist MD, Bigler J, Patterson RE, Kristal AR, Shattuck AL, Potter JD, White E (2002). Buccal cell DNA yield, quality and collection costs: comparison of methods for large-scale studies. Cancer, Epidemiology, Biomarkers & Prevention 11:1130-3.
- Shutler GG, Gagnon P, Verret G, Kalyn H, Korkosh S, Johnston E and Halverson J (1999). Removal of a PCR inhibitor and resolution of DNA STR types in mixed human-canine stains from a five year old case. *Journal of Forensic Sciences* 44(3):623-626.
- Tsai YL and Olson BH (1992). Rapid method for separation of bacterial DNA from humic substances in sediments for polymerase chain reaction. Applied and Environmental Microbiology 58(7):2292-2295.

Copyright protects this publication. However, Queensland Health has no objection to this material being reproduced with acknowledgment, except for commercial purposes. Permission to reproduce for commercial purposes should be sought from:

Managing Scientist DNA Analysis Forensic and Scientific Services PO Box 594, Archerfield QLD Australia 4108

Or by telephone (07) 3274 9169



Page 21 of 21



Batching Validation Samples ~ Blood

Sensitivity Experiment

20 Cotton Samples
1 "Blank
1 Positive Control
1 Negative Control
VALB20070511_01
Extracted: CI 12/6/07

20 Rayon Samples
I "Blank
I Positive Control
I Negative Control
VALB20070511_02
Extracted: CI 8/6/07

Inhibition Experiment

8 Indigo Samples
1 "Blank
8 Urea Samples
1 "Blank
1 Positive Control
1 Negative Control
VALB20070511_03
Extracted: CI 15/6/07

8 Humic Samples
1 "Blank 1 "Blank
8 Tannic Samples 1 Positive Control
1 "Blank 1 Negative Control
1 Positive Control
1 Negative Control
VALB20070511_05
Extracted: VH 8/6/07
Extracted: CI 14/6/07

Substrate Experiment

4 Denim Samples
1 "Blank
4 Cotton Shirt Samples
1 "Blank
4 Wool Samples
1 "Blank
4 Lycra Samples
1 "Blank
1 Positive Control
1 Negative Control
VALB20070511_06
Extracted: CI 13/6/07

4 Nylon Samples
1 "Blank
4 Polyester Samples
1 "Blank
1 Positive Control
1 Negative Control
VALB20070511_07
Extracted: CI 7/6/07

4 Rayon Swabs
1 "Blank
4 Cotton Swabs
1 "Blank
4 FTA Donor 1 Samples
4 FTA Donor 2 Samples
1 FTA Blank
1 Positive Control
1 Negative Control
VALB20070511_10
Extracted: GSL 15/6/07

4 Leather Samples
1 "Blank
1 Positive Control
1 Negative Control
VALB20070511_08
Extracted: CI 5/6/07

Mixture Experiment

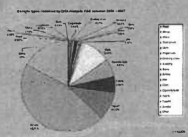
4 1:1 F:M Samples 4 1:2 F:M Samples 4 1:10 F:M Samples 1 " Blank 1 Positive Control 1 Negative Control VALB20070605_02 Extracted: GSL 13/6/07 4 1:25 F:M Samples 4 1:50 F:M Samples 4 1:100 F:M Samples 1 " Blank 1 Positive Control 1 Negative Control VALB20070605_03 Extracted: BG 14/6/07

Size Experiment

12 Cotton Shirt Samples
1 " " Blank
1 Positive Control
1 Negative Control
VALB20070511_11
Extracted: BG 12/6/07

Chemical Validation of a Manual DNA Extraction Method using the DNA IQTM System Jenny Lundie, Iman Muharam, Breana Gallagher, Cecilia lannuzzi, Vojtech Hlinka, Thomas Nurthen, Vanessa lentile

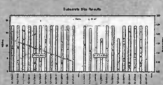
Abstract





Methods





Results and Discussion

Conclusions

1.8ased on overall results of the DNA IQ ** system validation, we concluded that the DNA IQ ** system is suitable for processing call and blood samples for DNA Analysis FSS.
2.The DNA IQ ** system can be verify on the MultiPROBE *II PLUS INT EX platforms for automated DNA extraction of various DNA Analysis FSS substrate types .

References

Acknowledgements

I wish to thank the the following for thair esistance and support

1. Australian and New Zestland Forensic Science Society. OLD Brench, Australia, 2. Promope Corporation/Med

6. ORA Analysis (55. CAMSCHE) Profiled Section, Australian and E. Mil Treatmentation Profiled to and OLD Australia.

CaSS Forensic and Scientific Services



CaSS Forensic and Scientific Services

Project 13. Report on the Verification of an Automated DNA IQ™ Protocol using the MultiPROBE® II PLUS HT EX with Gripper™ Integration Platform

Nurthen, T., Hlinka, V., Muharam, I., Gallagher, B., Lundie, G., lannuzzi, C., lentile, V. Automation/LIMS Implementation Project, DNA Analysis FSS (August 2008)

1. Abstract

A manual method for extracting DNA from forensic samples using the DNA IQ™ system (Promega Corp., Madison, WI, USA) was validated for routine use in DNA Analysis (FSS). We have verified an automated DNA IQ™ protocol in 96-well format for use on the MultiPROBE® II PLUS HT EX Forensic Workstation platforms (PerkinElmer, Downers Grove, IL, USA). Data indicate that results from the automated procedure are comparable to those from the manual procedure. Contamination checks were performed using samples prepared in checkerboard and zebra-stripe format, and results were as expected. We recommend the use of the MultiPROBE® II PLUS HT EX platforms to perform automated DNA extraction using the DNA IQ™ system.

Introduction

The MultiPROBE® II PLUS HT EX FORENSIC WORKSTATION platforms (PerkinElmer, Downers Grove, IL, USA) are equipped to perform automated DNA extractions, as they include a DPC shaker and individual heat controllers to enable on-board lysis and incubation steps. Currently in DNA Analysis, the MultiPROBE® platforms allow walk-away operation of PCR setup protocols for DNA quantitation and amplification.

The DNA IQ™ protocol has been verified or validated by various laboratories for use on the MultiPROBE® II PLUS platform. The laboratories that perform an automated DNA IQ™ protocol include PathWest (Western Australia), Forensic Science South Australia (South Australia) and Centre of Forensic Sciences in Toronto (Ontario). The MultiPROBE® II PLUS instrument comes pre-loaded with an automated DNA IQ™ protocol. Unlike the other boratories, however, we did not validate the included protocol, but instead validated a manual DNA IQ™ protocol which was based on the CFS automated protocol (PerkinElmer, 2004), followed by verification of an automated protocol based on the validated manual method.

The verified automated DNA IQ™ protocol is identical to the validated manual protocol used in-house: there are no differences in reagents or volumes. The adopted DNA IQ™ protocol differs slightly, however, from the manufacturer's protocol, as it includes a lysis step using Extraction Buffer (10mM Tris, 1mM EDTA, 100mM NaCl, 20% w/v SDS) in the presence of Proteinase K, before incubating in the DNA IQ™ Lysis Buffer. Furthermore, the lysis incubation conditions were lowered from 70°C to 37°C in order to accommodate extraction of DNA from heat labite materials such as nylon and polyester. In addition, the automated protocol utilises the SlicPrep™ 96 Device (Promega Corp., Madison, WI, USA) for simultaneous processing of samples in a 96-well format.



Page 1 of 18

CaSS | Forensic and Scientific Services

To verify an automated DNA IQ™ protocol for use on the MultiPROBE® II PLUS HT EX platforms to allow extraction of DNA from various sample types.

Equipment and Materials

- MultiPROBE® II PLUS HT EX with Gripper™ Integration Platform (PerkinElmer, Downers Grove, IL, USA)
- Gravimetric Performance Evaluation Option with Mettler SAG285/L balance (Mettler-Toledo,
- Greifensee, Switzerland) DNA IQ™ System (Promega Corp., Madison, WI, USA)
- Extraction Buffer (10mM Tris, 1mM EDTA, 100mM NaCl, 20% SDS)

- StilcPrep™ 96 Device (Promega Corp., Madison, WI, USA)
 Nunc™ Bank-It tubes (Nunc A/S, Roskilde, Denmark)
 175µL non-conductive sterile filter RoboRack tips (PerkinElmer, Downers Grove, IL, USA)
- 1000µL Conductive sterile filter Robotix tips (Molecular BioProducts, San Diego, CA, USA)
- ABI Prism® 7000 SDS (Applied Biosystems, Foster City, CA, USA)
- Quantifiler™ Human DNA Quantification kits (Applied Biosystems, Foster City, CA, USA)
- AmpFtSTR® Profiler Plus Amplification kits (Applied Biosystems, Foster City, CA, USA)
- GeneAmp® 9700 thermalcycers (Applied Biosystems, Foster City, CA, USA)
- ABI Prism® 96-well optical reaction plates (Applied Biosystems, Foster City, CA, USA)
- ABI Prism® 3100 Genetic Analyzer (Applied Biosystems, Foster City, CA, USA)
- GeneScan™ 500 ROX™ Size Standard (Applied Biosystems, Foster City, CA, USA)
- HI-Di™ Formamide (Applied Biosystems, Foster City, CA, USA) 3100 POP-4™ Polymer (Applied Biosystems, Foster City, CA, USA)
- Cytobrush® Plus Cell Collector (Cooper Surgical, Inc., Trumbull, CT, USA)
- 0.9% saline solution (Baxter Healthcare, Old Toongabbie, NSW, Australia)
- Stem digital tilting head thermometer
- For mock samples:
 - FTATM® Classic Card (Whatman Inc., Florham Park, NJ, USA)
 - Sterile cotton swabs (Medical Wire & Equipment, Corsham, Wiltshire, England)
 - Sterile rayon swabs (Copan Italia SPA, Brescia, Italy)

Methods

5.1 Gravimetric Evaluation of Pipetting Accuracy and Precision

Gravimetric analysis was performed by placing the SAG285/L balance on the platform deck and instructing the MP II to repeatedly pipette certain volumes of system liquid onto the balance pan. Readings were taken automatically by the software and compiled into a results table, which was then used to automatically generate an Excel-based results chart containing mean, %CV and %inaccuracy values. The mean values obtained were used to calculate R2, slope and Y-intercept (offset) values to calibrate the system's pipetting.

Pipetting performance was assessed for various volumes using three different tips in order to calculate appropriate R2, slope and Y-intercept (offset) values which were then added to the performance file. Values were calculated for both Blowout (single-liquid transfer) and Waste (multidispense) modes for the 1000µL conductive tips, and Blowout mode only for the 175µL non-conductive tips and fixed tips.

For the addition of resin, a specialised performance file was created based on the performance file for 175µL tips in blowout mode, except the "Blowout Volume" column values were set to 0 to allow pipetting performance that is similar to waste mode. Retesting was performed to confirm accurate and precise pipetting with these settings.



Page 2 of 18

FSS.0001.0084.1446

CaSS Forensic and Scientific Services

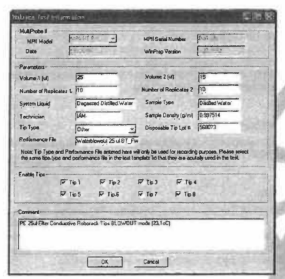


Figure (SEQ Figure * ARABIC). The Balance Test Information Window as present within the Balance Test DT program. All pipetting parameters are entered here and are subsequently transferred to the result output file.

All gravimetric testing was performed using the Balance Test DT test program within WinPrep®. Parameter values that needed to be entered into the Balance Test Information Window (Figure 1) included those as outlined in Table 1.

Table { SEQ Table * ARABIC }. Input values that are required for the various Balance Test Information parameters.

Parameter(s)	Value
Volume 1 and Volume 2	For 175µL tips: 175, 100, 50, 15µL
	For 1000µL tips: 1000, 700, 400, 100µL
	For fixed tips: 1000, 700, 400, 100µL
Number of Replicates	10
System Liquid	Degassed Nanopure Water
Sample Type	Nanopure Water
Technician	Initials of the operator performing the test
Sample Density (g/ml)	The density of water at environmental temperature*
Tip Type	Other
Disposable Tip Lot #	The lot number of the particular tips in use
Performance File	The appropriate Performance File for the tip (175µL, 1000µL or fixed
	tips) and pipetting mode (Blowout or Waste) in use
Enable Tips (checkboxes)	Select the actual tips (1 to 8) to be tested
Comments	Free text box to add additional information (eg. Tip type, mode, current environmental room temperature, etc).

^{*} Water density values were obtained from http://www.simetric.co.uk/si_water.htm

Pipetting accuracy and precision were examined at four different volumes for each tip size: 175, 100, 50, 15 μ L for 175 μ L tips and 1000, 700, 400, 100 μ L for the 1000 μ L and fixed tips.



Page 3 of 18

CaSS | Forensic and Scientific Services

In order to calculate unbiased values for each set of volumes, the slope and offset values in the relevant Performance File were changed to the default 1 and 0 respectively prior to testing. The mean volumes that were pipetted by each tip (10 replicates per tip) at the four designated volumes were used to generate a standard curve. The slope and offset calculated from this curve was used to calibrate the relevant Performance File. The final Performance File settings were then tested at the highest and lowest volumes (as per Table 1) to confirm accurate and precise pipetting.

5.2 Blood Collection

Blood samples were collected from 2 staff donors (DJC/VKI) by a philebotomist as per normal in three 4mL EDTA vials. Blood samples were stored at 4°C.

5.3 Cell Collection

Buccal cells were collected using a modified Cytobrush® protocol (Mulot et al., 2005; Satia-Abouta et al., 2002). The donor was instructed to brush the inside of one cheek for one minute using a Cytobrush®. Then, with another Cytobrush®, the other cheek was also sampled. Once each cheek was swabbed, the cells on the brush were suspended in 2mL of 0.9% saline solution. Buccal cell samples were stored at 4°C.

5.4 FTA cell Collection

Cells were collected from two staff donors (VKI/CJA) by using a "lolly-pop" swab to sample the inside of the donor's cheek for 15 seconds before pressing the swab onto the FTA™ paper to transfer the DNA. FTA™ cards were stored at room temperature.

5.5 Heater tile temperature verification

Heat tiles supplied with the MultiPROBE® II PLUS HT EX platforms were modified to accept the SlicPrep™ 96 Device. For testing, 1mL of nanopure water (at room temperature) was added to each well. The plate was then placed on a heater tile (controlled by the MP II heater controller) and allowed to reach temperature. The temperatures tested were 37°C and 65°C. Temperature readings for specific outer and inner wells (i.e. A1, A6, A12, D1, D6, D12, H1, H6, H12) were taken at regular intervals up to and including 45 minutes, using calibrated stem digital tilted head thermometer probes. The data were collated and means calculated to determine the distribution of heat over the tile

5.6 Verification of automated DNA IQ™ Protocol

The automated DNA IQ™ protocol, based on the validated manual method (refer to Project 11), was programmed in WinPrep™ software. The final, optimised protocol was named "DNA IQ Extraction_Ver1.1.mpt". A screenshot of the Test Outline window for this protocol is depicted below in Figure 2. The deck layout is illustrated in Figure 3.

The automated DNA IQ™ protocol was designed to mimic the validated manual method, with minor modifications. Briefly, the changes include:

- o Increasing the volume of Extraction Buffer to 500µL;
- A SlicPrep™ 96 Device (Promega Corp., Madison, WI, USA) was used for sample lysic:
- Incubation steps and any shaking steps were performed on the integrated DPC shaker:



Page 4 of 18

FSS.0001.0084.1448

CaSS | Forensic and Scientific Services

- o CRS toroid magnet (P/N 5083175) was used for isolating the DNA IQ™ resin.
- o Instead of a single elution of 100μL, a double elution method (2 x 50μL) is used.

Reagents used in the automated protocol were as per the manual method.

```
Test Outline
    Test Outline

2 Initial User Query (x 1)

3 L.-BercodeSetup (x 1)

1 3 User Nessage - Hardware setup (x 1)

3 User Nessage - Hardware setup (x 1)

5 5 OberComm to Shaker (x 1)

6 5 Set, Heater Temperature at 37 C (x 1)

7 5 6 Set, Heater Temperature at 65C (x 1)

8 7 6 Add SOUL Extraction Buffer to SicBask (x File: Records)
                                                                                                                                                   ( x 1)
                                                                                                                                              52. Shaker On Wash 2 (x 1)
                                                                                                                                                   1 54. StopShakerWash2 (x 1)
                                                                                                                                              6. 75. Flosh WashWash2 ( x 1.)
6. Move Plate SlicPrep to PKI.MagnetWash2 ( x 1.)
                                                                                                                                              57. Bind 1 minute_Wash2 (× 1.):
58. Remove wash buffer 2 ( x.File: Records)
                                                                                                                                              9. Wait for 37 Temperature ( x 1.)
10. Seal plate ( x 1.)
     11. ShakerOn_L(x1)

2 42. Incubate 45 min on heater/shaker_I (x1)
                                                                                                                                              . 3. 64. StopShakerWash3 (x 1 )

3. 65. Flush(WashWash3 (x 1 )

3. 14 66. Move Place SlicPrep to PSI MagnetWash3 (x 1 )
     13. StopShaker_1 (x l.)
14. Centrifuge (x l.)
15. Place SlicPrep D16 (x l.)
                                                                                                                                             ■ $ 6.8 Bind 1 minute_Wash3 (x.1.)

■ $ 6.8 Remove wash buffer 3 (x.File: Records )

■ $ 69. Dry 5 minutes (x.1.)

■ $ 70. Flush (Wash_4 (x.1.)
         16: Flush (Wash_1 (x l)
12: 17. Add Resin 50uL (x File: Records)
     ** $8. Flush Mash 3 (x 1)

** 19: Add DNA IQ Lysis Buffer (957. ul) to SlicPrep at 016 (x File: Records)

** 20. Flush Mash 1 (x 1)

** 40. 21. Move Plate_1 (x 1)
                                                                                                                                              71. Wait for 65 Temperature_1 (x 1)
72. Add Budion Buffer (60ut) Eluti (x File: Records )
73. Move Shorrep from PK1 Magnet to Tiez on Shaker_1 (x 1)
         22, ShakerOn, 2 (x 1)
23. Time 5 min_1 (x 1)
                                                                                                                                              9- X 74.3 minutes Timer_1 (x1)
                                                                                                                                             75. ShakerOnElut I.(x1)
     24. StopShaker_2(x1)

25. Move SlicPrep to PKI Magnet (x1)
                                                                                                                                             77: StopShakerElu1 ( x 1 ).

78: 78: Move SlicPrep from Tile2 to PKI:Magnet_1 ( x 1 ).
    ②- 

26: Time 1 mln.- Walt to Bind Resin_1 (x.1)

27: 27. Remove 1600ut to AxSuper (x File: Records)
                                                                                                                                             # 79. Push Down-Skripe Elut 1 (x 1)

80. Bind 1 minute Elut 1 (x 1)

81. Transfer Eluted DNA_Elut 1 (x File: Records)
     30. Dispense Lysis Buffer (125 ul) ( x File: Records )
                                                                                                                                             31. Flush/Wash_4(x1)
    32, $hakerOn_3(x1)

33, Timer_1(x1)

34, $top$haker_3(x1)
                                                                                                                                              3 14 84: Move SlicPrep from PKI Magnet to Tile2 on Shaker_2 ( x 1 )

        ■ 85. 3-minutes Timer_2(x1)

        ● 86. 5-bakerOnElut2(x1)

        ● 2 87. 5-bake 3 minute Elut2(x1)

    35. Flush/Wash_1 ( x 1 )

36. Move SlicPres to PKI Magnet ( x 1.)
                                                                                                                                              88. StopShakerElu2 ( x 1 )
    37, Time 1 minute ( x 1 ) 38. Remove Lysis Buffer (125 ul) to STORE ( x File: Records )
                                                                                                                                            ## 88. StopShakerELUZ (x 1)
## 189. Move SlicPrep from Tile2 to PKI Magnet_2(x 1)
## 190. Push Down SlePrep Edut2(x 1)
## 191. Bind 1 minuterEkit2(x 1)
## 192. Transfer Eluted DNA_Elut2(x File: Records)
## 193. Flush/Wash_6 (x 1)
## 194. Gose Heater Comm(x 1)
## 195. Gose Shaker Comm(x 1)
## 196. Remove Nunctubes (x 1)
## 197. Amphyd concentrate (x 8)
## 198. Amphyd dilute (x 8)
    39. Move SicPrep from PKI Magnet to Shaker 1 ( x 1:)
40. Add wash buffer 1 ( x File: Records )
    41. Flush/Wash_1(x1)
42. ShakerOnWash1(x1)
    43. Shake 1 minute.Wash! (×1)
    15. Flush/WashWash1 ( x 1 )
                                                                                                                                             E- 73 98. Amphyl. dilute ( x 8 )
    选 X 47. Bind 1 minute_Wash1 (×1)
                                                                                                                                             1 72 99. Water wash (XB)

    48. Remove wash buffer I (x File: Records)
    49. Move SlicPrep from PKI Magnet to Shaker 2 (x 1)
    50. Add wash buffer 2 (x File: Records)
                                                                                                                                             100, Flush/Wash_5 ( x 2 )
                                                                                                                                                   O End of Test
```



Page 5 of 18

CaSS | Forensic and Scientific Services

Figure 2. The Test Outline window displaying individual nodes within the DNA IQ Extraction_Ver1.1.mpt program test file

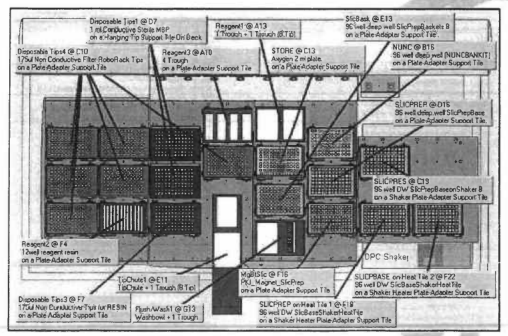


Figure 3. The deck layout for DNA IQ Extraction_Ver1.1.mpt, displaying the required labware on the platform deck.

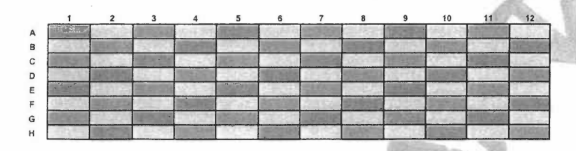
The automated DNA IQ™ protocol was used to perform the following tests.

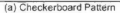
5.6.1. Contamination Check via Checkerboard and Zebra-stripe Patterns Samples consisting of two 3.2mm FTA® discs (containing blood, buccal cells, or blank cards) were arranged in a checkerboard and zebra-stripe pattern (Figure 4) in SlicPrep™ plates using the BSD Duet 600 instrument (BSD Robotics, Brisbane, QLD, Australia) and extracted on the MultiPROBE® II PLUS HT EX platforms using the automated DNA IQ™ protocol. One checkerboard and one zebra-stripe plate was processed on each platform.



Page 6 of 18

CaSS | Forensic and Scientific Services





b) Zebra Stripe Pattern

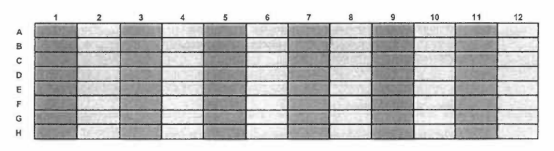




Figure 4. Checkerboard and zebra-stripe patterns utilised in the contamination check.

5.6.2. Comparisons with the manual DNA IQ™ method

Comparisons were made between results generated by the automated and manual methods to verify the performance of the automated DNA IQ^{TM} protocol.

Verification samples consisted of different dilutions of blood and cells spotted in 30µL aliquots onto quartered cotton and rayon swabs. Four blood dilutions of neat, 1/10, 1/100 and 1/1000 and four cell dilutions of neat, 1/5.2, 1/52.2 and 1/522 were used to test the sensitivity of both the manual and automated methods. Dilutions were created using 0.9% saline solution for both sample types. Four replicates of each dilution were made up for each substrate and sample type.



Page 7 of 18

CaSS | Forensic and Scientific Services

The blood was collected using the same method as in 5.2. Four separate extractions were performed for the manual set based on the combination of sample type and swab type: Blood Rayon, Blood Cotton, Cell Rayon and Cell Cotton. For the automated verification, all sample types were extracted together after being transferred to a SlicPrep™ 96 Device to allow automated processing.

5.6.3. Resin volume

The performance of the automated DNA IQ™ protocol was assessed when either 7µL or 14µL of DNA IQ™ resin was used in the protocol to extract blood samples.

5.6.4. Modifying extraction volumes

The performance of the automated DNA IQTM protocol was assessed for varying volumes of extraction buffer at 300, 350, 400, 450 and $500\mu L$. In each case, the volume of DNA IQTM Lysis Buffer was kept at 2x the volume of extraction buffer. Samples extracted were blood swabs, prepared as

5.6.5. Sensitivity of the automated DNA IQ™ protocol

The sensitivity of the automated DNA IQ™ protocol was assessed using dilutions of whole blood at neat, 1:10, 1:50, 1:100 and 1:1000.

6. Results and Discussion

6.1 Gravimetric Evaluation of Pipetting Accuracy and Precision

Pipetting on both automated platforms was assessed gravimetrically as per laboratory practice. Gravimetric results indicate that pipetting performance for five different pipetting behaviours using $500\mu L$ syringes on the instruments is accurate and precise to within the established threshold of $\pm 5\%$ (Table 2). The maximum CV at the maximum volume was 0.78%, whereas the maximum CV at the minimum volume was 1.1%. The CV for pipetting at lower volumes is expected to be slightly higher than the CV at higher volumes using $500\mu L$ syringes, because accuracy at small volumes is harder to achieve with larger syringe sizes. Nevertheless, pipetting on the extraction platforms is limited to a minimum of $50\mu L$, which exhibited a CV of 0.36%.

Table 2. Gravimetric evaluation results for various performance files used on either MP II EXTN A or MP II EXTN B.

Performance File	Max. Vol. µL	Min. Vol. µL	Max. Vol. µL Mean	Max. Vol. %CV	Max. Vol. %Inac.	Min. Vol. µŁ Mean	Min. Vol. %CV	Min, Vol. %Inac.
EXTN A							120	
Water Blowout 175µL DT_FW _13112007RESIN.prf	50µL	N/A	49.98	0.36	0.0	N/A	N/A	N/A
Water Blowout 175µL DT_FW QHSS_13112007.prf	175uL	15µL	172.26	0.21	1.6	425.47	STEEN :	HENR
WaterWaste 1mL_FW_QHSS 12112007.prf	1000µL	100µL	999.11	0.24	0.1	99.22.	0.71	0.8
Water.Blowout 1mL DT_QHSS_09112007.prf	1000µL	100µL	1001.02	0.27	0.1	100.65	0.63	0.7
Water Blowout Fixed Tips_08112007.prf	1000µL	100µL	995.97	0.31	0.4	99.6	0.71	0.4
EXTN B								
Water Blowout 175µL DT FW 25102007RESIN.prf	50uL	N/A	50.12	0.36	0.2	N/A	N/A	N/A
Water Blowout 175µL DT_FW_25102007.prf	175µL	15µL	175.58	0.14	0.3	15.23	1.1	1.5
WaterWaste 1mLDT_FW_QHSS 24102007.prf	1000uL	100µL	1002.39	0.78	0.2	99.56	0.89	0.4
Water Blowout 1mL DT QHSS 23102007.prf	1000uL	100µL	998.2	0.44	0.2	99.44	0.68	0.6



Page 8 of 18

CaSS | Forensic and Scientific Services

Water Blowout Fixed Tips FW 26102007.pdf	1000uL	100uL	998.87	0.68	0.1	100.37	0.74	0.4

6.2 Heater tile temperature verification

Two heater tiles on each MP II platform was verified to reach either 37°C or 65°C, the optimum incubation temperatures for sample lysis and DNA elution respectively (using the DNA IQ™ kit). Each tile, upon completion of the verification process, could only be used for a specific temperature, and as such was labelled appropriately to ensure use of the correct tile for specific incubation steps (Table 3).

Table 3. Verified heater tiles for use in the automated DNA IQ™ protocol.

Extraction platform	Tile number	Heater Controller Setting	Average °C reached	Verified temperature	Incubation Step
EXTN A	3 (45W)	50°C	37°C	37°C	Sample Lysis
EXTN A	1 (45W)	85°C		65°C	DNA Elution
EXTN B	1 (45W)	50°C		37°C	Sample Lysis
EXTN B	2 (45W)	85°C	65°C	65°C	DNA Elution

A slight variation in the incubation temperature to achieve sample lysis is acceptable, because Proteinase K exhibits stable activity and broad specificity over a wide range of temperatures between 20-60°C, at which the serine protease still retains greater than 80% of its activity (Sweeney & Walker, 1993).

The efficiency of the elution step is dependent on heating the sample to 65°C in the presence of DNA IQ™ Elution Buffer (Huston, 2002). If the sample is not sufficiently heated, the extraction yield may be lower than expected. Two heater tiles were able to be verified for this crucial incubation step, with both tiles exhibiting minimal variation.

6.3 Contamination Check via Checkerboard and Zebra-stripe Patterns
Table 4 below lists the Extraction Batch ID's of the contamination checks.

Table 4. Extraction Batch ID's for the various contamination check plates that were processed on the MP II platforms using the automated DNA IQ^{TM} protocol.

Type of plate	Extraction batch ld	Extraction Platform	Check passed
Checkerboard 1	VALB20070817_02	Extraction A	Invalidated
Checkerboard 2	VALB20070803 02	Extraction B	Yes
Zebra-Stripe 1	VALB20070803_03	Extraction A	Yes
Zebra-Stripe 2	VALB20070817_03	Extraction B	Yes
Checkerboard/Zebra	VALB20071022 01	Extraction A	Yes

Checkerboard 1

Position E3 Sample Cells 6) was known to have been contaminated prior to the start of the extraction. The result showed a mixed DNA profile, with contributing alleles originating from the expected wells (Table 5). In addition to this contamination event, eight of the designated blank samples (positions D3, A10, F1, H5, C4, E4, B7 and E6), two of the cell samples (A1 and B10) and two of the blood samples (F4



Page 9 of 18

FSS.0001.0084.1453

CaSS Forensic and Scientific Services

and G7) all exhibited a partial DNA profile that was previously unknown (Table 5). This profile did not match any of the positive control samples present on the batch. The DNA profile was searched against the Staff Database and no matches were found. The source of this contaminating DNA profile could not be identified.

None of the other blank samples yielded any DNA profile. The rest of the cell and blood samples yielded the correct DNA profile. Although there is no evidence of well-to-well contamination, the unknown DNA profile obtained has invalidated this plate. A further checkerboard/Zebra-Stripe combination plate was performed to ensure...

Table 5. The DNA profile of the unknown contaminant that was observed in Checkboard-1.

Sample description	D3	vWA	FGA	Amel	D8	D21	D18	D5	D13	D7
Blk23-E6	14,17	14,17	22,24	X,Y	11,11	29,32.2	14,15	9.11	11,12	11,13
Blk25-B7	14,17	14,17	22,24	X,Y	11,11	29,32.2	14,15	9,11	11,12	
Blk15-E4	14,17	14,17	22,24	X,Y	11,11	29.32.2	14.15	9,11	11,12	11,13
Blk14-C4		14		X	11	32.2	TOLDS	9		
Blk20-H5	14,14	17,17	20,21	X.X	13,16	29.30	14,16	11,13	11,12	11,11
Blk3-F1	14	17		X	13	29,30	14	CONT.	12	11
Blk10-D3	14,17	14		X,Y	11	29,32.2	14	9,11		11,13
Blk37-A10	14,17	14	22,24	X,Y	11	29	14	9,11		
Cells19- B10	14,17	14,17	20,21,22,24	X,Y	11,13,16	29,30,	14,15,16	11,15	11,12	11,11
Cells13-A1	14.17	14,17	20,21,22,24	X,Y	11,13,16	29,30,32,32,2,33	14,15,16	9,11,13	11,12	11.13
Blood14-G7	NR.17,18	NR.16	20,21	X.Y	NR.13,14	29,30,31,NR	NR.14	NR.12	10,10	10,NR,12
Blood8-F4	NR, 17, 18	NR, 16, 17	20,21,NR,24	X,Y	11,13,14	29,30,NR,NR	14,14	9,11,12	10,NR	10,NR,12
Cells 6-E3	14,17,18	16,17	20,21	X,Y	13,14,16	29,30,31	NR,14,16	11,12,13	10,11,12	10,11,12

Checkerboard 2

None of the blank samples yielded DNA profiles; all of the positive cell and positive blood samples yielded the correct DNA profile. Figure 5 illustrates the DNA quantitation results from this plate. DNA was not detected in any of the blank samples.



Page 10 of 18

CaSS Forensic and Scientific Services

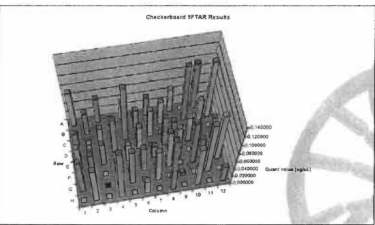


Figure 5. Checkerboard 2 quantitation results, showing the absence of detectable DNA in the blank samples (grey).

Zebra-Stripe 1

None of the blank samples yielded DNA profiles, all of the positive cell and positive blood samples yielded the correct DNA profile. Figure 6 illustrates the absence of detectable DNA in the blank samples.

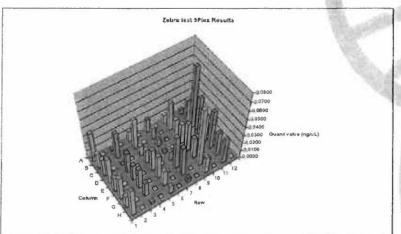


Figure 6. Zebra-Stripe 1 quantitation results, showing the absence of detectable DNA in the blank samples (grey).

Zebra-Stripe 2

None of the blank samples yielded DNA profiles, all of the positive cell and positive blood samples yielded the correct DNA profile. Figure 7 shows the absence of detectable DNA in the blank samples.



Page 11 of 18

FSS.0001.0084.1455

CaSS | Forensic and Scientific Services

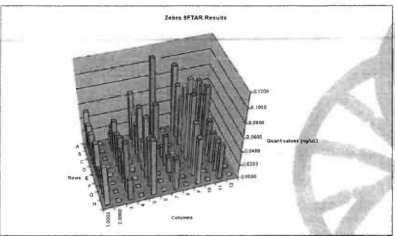


Figure 7. Zebra-Stripe 2 quantitation results, with no DNA detected in the blank samples.

Checkerboard/Zebra

None of the blank samples yielded DNA profiles, all of the positive cell and positive blood samples yielded the correct DNA profile. DNA was undetected in the blank samples (Figure 8).

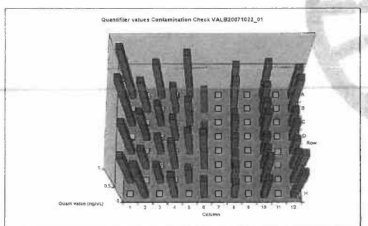


Figure 8. Checkerboard/zebra plate that was extracted on MP II Extraction Platform A because the previous plate was invalidated. DNA was not detected in the blank samples (grey).

6.4 Comparisons with the manual DNA IQ™ method

When dilutions of either blood or cells were applied on to either rayon or cotton swabs, followed by extraction using the DNA IQ™ method, the results of the automated method were always lower in yield compared to the manual method. For blood samples on rayon



Page 12 of 18

FSS 0001 0084 1456

CaSS | Forensic and Scientific Services

swabs, the automated method generated yields that were on average around 8% (SD 8.45%) of the automated method. For blood on cotton swabs, the yield from the automated method was also around 8% (SD 3.62%). The yields for cell samples were higher at around 33% (SD 16.29%) and 25% (10.32%) for cells on rayon and cotton swabs respectively.

The manual method was found to be more sensitive than the automated method. Out of five replicates at the 1/100 and 1/1000 dilutions for blood on rayon swabs that were processed using the manual method, five and three replicates respectively were detected (and none from the automated method) (see Figure 9). The trend is repeated for blood on cotton swabs (Figure 10). For cell samples on either rayon or cotton swabs, the automated method was found to be more sensitive as evidenced by detection of DNA at the 1/522 dilutions (Figure 11 and 12).

Cell clumping may have occurred with the cell dilutions, therefore causing inaccurate dilutions as can be observed in the ratios between each dilution.

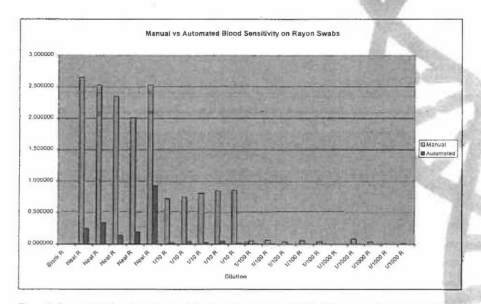


Figure 9. Comparison of sensitivity between the manual and automated DNA IQ™ methods for blood samples on rayon swabs.



Page 13 of 18

CaSS | Forensic and Scientific Services

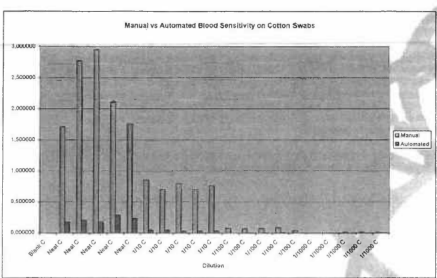


Figure 10. Comparison of sensitivity between the manual and automated DNA IQ™ methods for blood samples on cotton swahs

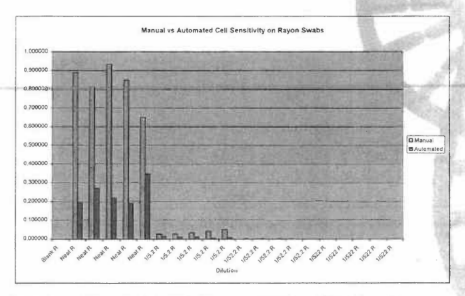


Figure 11. Comparison of sensitivity between the manual and automated DNA IQ^{TM} methods for cells samples on rayon swabs.



Page 14 of 18

CaSS | Forensic and Scientific Services

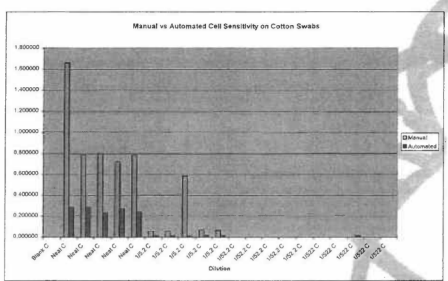


Figure 12. Comparison of sensitivity between the manual and automated DNA IQ™ methods for cell samples on cotton swabs.

6.5 Investigating resin volume

Promega recommends the use of $7\mu L$ of DNA IQ^{TM} resin with their protocol. We investigated the performance of the protocol with double the amount of resin $(14\mu L)$ in order to assess any benefits that may be gained in terms of the resulting yield and quality of the STR profile.

It was observed that doubling the resin resulted in a proportional doubling of the yield. On average, doubling the resin increased the yield by an additional 77.28% (n=4). The average yield from an extraction using 7 μ L of resin was 64.725ng (SD 32.21ng, n=4), whereas 14 μ L resin generated 114.75ng (SD 10.72ng, n=4) (Table 6). At the higher resin concentration, the amount of DNA isolated appears to be capped at around 100ng, indicating no change in the ability of the reaction to isolate more DNA due to saturation of resin.

Table 6. Comparison of the effects of doubling the amount of

ecommended DNA IQ™ resin.

Sample ID	Resin volume	(DNA) ng/µL	Reportable alleles
33383-4216		0.701	18/18
33383-4225	71.1	1.070	18/18
33383-4239	7µL	0.319	18/18
33383-4248		0.499	18/18
33383-4252		1.140	18/18
33383-4261	4.41	1.270	18/18
33383-4270	14µL	1.010	18/18
33383-4284		1.170	18/18



Page 15 of 18

CaSS Forensic and Scientific Services

Samples extracted using either amount of resin generated concordant full DNA profiles (18/18 alleles). Samples processed using the 14µL method produced peaks that were slightly higher. The difference in peak heights between alleles within the same loci ranged from 59-86%, with a mean of 71%, indicating minimal difference between the two methods.

Doubling the amount of resin did not appear to provide any additional benefits compared to the original recommended protocol. More importantly, full DNA profiles were resolved using either method. Therefore, the costs associated with increasing the amount of resin cannot be justified at this stage.

6.6 Modifying extraction volumes

An investigation into optimising extraction volumes ranging from 300µL to 500µL was performed in order to ensure that buffer coverage over the samples was sufficient to enable optimal lysis and release of DNA. In addition, the use of an optimum volume of extraction reagents increases efficiency and economy, therefore potentially lowering laboratory costs.

Although the higher extraction volume generated higher yields when processed using the automated DNA IQ™ protocol (Table 7), DNA profile results were comparable across the various extraction volumes tested for eight replicates each (Table 8). Three instances of allelic imbalance were encountered in two samples from the 300µL and 450µL tests. In all instances, allelic imbalance was greater than 69%.

Table 7. DNA profile results for samples extracted using various volumes of Extraction Buffer, for 8 reglicates.

Extraction Buffer Volume (µL)	Mean [DNA] (ng/µL)	SD
300	2.04	0.07
350	2.16	0.09
400	1.69	0.10
450	3.14	0.13
500	3.64	0.17

Table 8. DNA profile results for samples extracted using various volumes of Extraction Ruffer, for 8 replicates

Sample	Extraction Buffer Volume (µL)	DNA Profile Result	
300-1 swab		OK	
300-2 swab		OK	
300-3 swab		OK	
300-4 swab	300	OK	
300-5 swab	300	OK	
300-6 swab		OK	
300-7 swab		AI D13	
300-8 swab		OK	



Page 16 of 18

CaSS Forensic and Scientific Services

350-1 swab 350-2 swab 350-3 swab 350-4 swab 350-5 swab 350 350-5 swab 350-6 swab 350-7 swab 350-8 swab 400-1 swab 400-2 swab 400-3 swab 400-4 swab 400 400-5 swab 400-6 swab 400-7 swab 400-8 swab 450-1 swab 450-2 swab OK 450-3 swab 450-4 swab 450-5 swab 450 450-6 swab 450-7 swab 450-8 swab 500-1 swab 500-2 swab

500

OK OK OK

6.7 Sensitivity of the automated DNA IQ™ protocol

500-3 swab 500-4 swab 500-5 swab

500-6 swab 500-7 swab 500-8 swab

DNA was detected from samples that were diluted down to 1:1000 (Figure 13).



Page 17 of 18

CaSS Forensic and Scientific Services

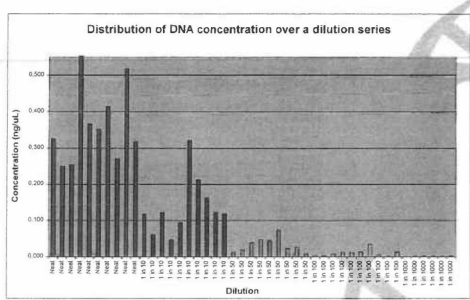


Figure 13. DNA IQ™ sensitivity across various dilutions

as

7. Summary and Recommendations

We recommend the following:

- Use of MPII for automated extraction of reference samples
- Use of MPII for automated extraction of casework samples
- Ongoing development of the automated extraction program to increase the efficiency of the extraction

Sweeney, P.J. and Walker, J.M., Burrell, M.M., Enzymes of molecular biology. *Methods Mol. Biol.* Towanam NJ, (1993) **16**, 306



Page 18 of 18