The Determination of Factors Involved in Column-Based Nucleic Acid Extraction and Purification

Jun-Jie Poh1* and Samuel Ken-En Gan2,3*

1Bioinformatics Institute, Agency for Science, Technology, and Research (A*STAR), 138671, Singapore
2p53 Laboratory, Agency for Science, Technology, and Research (A*STAR), 138648, Singapore
3Quintech Life Sciences Pte Ltd, 619933, Singapore

Abstract

DNA extraction methods such as plasmid minipreps, gel, and PCR purifications, are indispensable techniques for genetic manipulations. There are numerous factors that contribute to the efficiency of these processes, which determine the success of complex downstream molecular analytics and diagnostic tests. To study and optimize these factors, we compared our own proprietary buffers to commercially available column-based kits, utilizing their spin columns and protocols. Through systematic substitution of the buffers in the kits with our own proprietary buffers, we selected the highest DNA yielding buffer recipes. Further analysis of the differences between the buffers showed that high concentrations and presence of certain chaotropic agents and cations are necessary for good plasmid miniprep, gel extraction, and PCR purification kits.

Keywords: DNA extraction; Gel extraction; PCR purification; Column-based purifications

Abbreviations: E. coli; Escherichia coli; SDS: Sodium Dodecyl Sulfate; TE: Tris-EDTA; OPT: Optimized; HM: Home-Made

Introduction

The extraction and purification of nucleic acids are commonly used techniques to isolate genetic material from tissues, bacteria, plants, and viruses for important analytical, diagnostic and preparative downstream processes. Amongst these methods, plasmid DNA extraction was the first to be reported [1] using the tedious alkaline extraction protocol. This involved lysozyme treatment to weaken the Escherichia coli (E.coli) cell wall prior to cell lysis and selective denaturation of genomic DNA using sodium dodecyl sulfate (SDS) and sodium hydroxide. Sodium acetate is then used to neutralize the alkaline pH, resulting in the formation of an insoluble network of denatured genomic DNA, protein-SDS complexes and high molecular weight RNA. These complexes were then removed by high speed centrifugation, leaving the desired plasmid DNA in the supernatant [1].

As the protocol was labour-intensive, efforts to simplify the extraction methods gave rise to the development of the "Guainidinium Thiocyanate - Phenol - Chloroform" method [2] to separate the various biomolecules through multiple liquid phases [3]. Further developments resulted in doing away with the use of hazardous chemicals (phenol and chloroform) through the use of spin columns for rapid extraction of high purity nucleic acids. Despite simplifying the process through the immobilization of plasmid DNA to the solid phase matrix (i.e. silica), plasmid extraction is still underlined by the need to disrupt bacterial cell walls, denaturation of nucleic acid binding proteins, inactivation of nucleases such as RNases, washing away of undesired contaminants, and elution of desired plasmid DNA.

At the crux, the silica solid phase matrix determines the resultant product purity and yield. For optimal DNA binding, equilibration of these silica columns by Na+ is required to break hydrogen bonds for the formation of salt bridges, allowing for spatial interaction with the negatively charged DNA. The silica membrane is then washed with ethanol to remove salts and other contaminants prior to elution of the bound DNA using low ionic strength (pH ≥ 7) buffers [3].

Based on the same principle for plasmid DNA extraction, spin columns had also been used for gel extractions and polymerase chain reaction (PCR) purifications. While these developments have contributed greatly to biomedical research, major developments of these kits are generally largely focussed on membrane material science. To complement this, we have decided to investigate whether the manipulation of chemicals in associated buffers will increase the yields that would enable researchers to tweak their existing commercial kits for improved yields.

Materials and Methods

Investigation of miniprep buffers

**Proprietary buffers**: Proprietary equilibration buffers (P-BK1 and P-BK2); resuspension buffers (P-P1); lysis buffers (P-P2); neutralization buffers (P-P3-1 and P-P3-2); binding buffer (P-W1); wash buffers (P-W2-1 and P-W2-2) and elution buffers (P-EB1, P-EB2, and P-EB3) for plasmid extraction with the following ingredients were prepared:

| Buffer | Constituents |
|--------|--------------|
| P-BK1  | NaCl, MOPS   |
| P-BK2  | NaOH (> 1M [Na+] than P-BK1) |
| P-P1   | Tris Base, EDTA, RNase A |
| P-P2   | SDS, NaOH    |
| P-P3-1 | C2H5KO2      |
| P-P3-2 | NH4C(=NH)NH2 • HCl, C2H5KO2 (pH< P-P3-1) |

*Corresponding author: Samuel Ken-En Gan, Bioinformatics Institute, Agency for Science, Technology, and Research (A*STAR), 30 Biopolis Street, #07-01 Matrix, 138671, Singapore, Tel: 65-6478-8417, 65-6407-0584; Fax: 65-6478-9047; E-mail: samuel@g@bii.a-star.edu.sg

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Comparison of HM and OPT buffers on generic A and B plasmid extraction kits were each carried out in triplicates. DNA concentrations and A260/280 ratio were analysed spectrophotometrically using IMPLEN Nanophotometer P330 in triplicates. 10 µl of extracted/purified DNA from the above comparisons were loaded with 6x loading dye containing SYBR Green (Quintech Life Sciences) and analyzed using the RunVIEW electrophoresis apparatus (Cleaver Scientific). A and B gel extraction kits and proprietary buffers: Comparisons between gel extractions buffers from generic A, generic B, and the proprietary buffers (P-QG2, P-QG3) were performed in triplicates. Gel protocols for generic A and B (see Supplementary Material) were carried out according to respective manufacturer’s instructions with the exception of standardizing gel dissolution temperature to 60°C and elution of DNA at 35 µl. Gel extractions using proprietary buffers were carried out according to generic A protocol with the exception of varying the ratio of buffer to gel slice to 3:1 w/v ratio (according to generic B protocol). Time taken for the gel slices to dissolve completely were measured with a lab timer and analyzed statistically.

Optimization of PCR purification buffers

Polymerase chain reaction: PCR reactions of 325 µl were performed containing 6.5µl of Taq polymerase, 19.5 µl 30 mM MgCl2, and 32.5 µl of 10X PCR Buffer (Axil Scientific), 26 µl of 2 mM dNTPs mix (Quintech Life Sciences), 13 µl of reverse primer : OriP Nrul R (5’-ATA TCT CGC GAA TGC TGG GGG ACA TGTACC TC-3’), forward primer OriP Nrul F (5’-CAC ACT CGC GAA GGA AAA GGA CAA GCA GCG AA-3’), template plasmid DNA, and 201.5 µl of HyClone water (Thermo Scientific, Cat no. SH30383.01). The amplicon oriP is ~1.9 kb. The completed PCR mix was transferred into PCR tubes of 25 µl aliquots and carried out in Artik Theral Cyclcer (Thermo Scientific) with the following profile: Initial denaturation at 94°C for 5 minutes; 30 cycles of denaturation at 94°C for 1 minute, annealing and extension at 71°C for 3 minutes; and final extension at 72°C for 10 minutes.

Comparisons of PCR purifications of generic A and B, and proprietary buffers: Generic A and B PCR purifications and selected proprietary buffers from miniprep (P-W1) and gel extraction (P-QG2) buffers were carried out in triplicates using the respective generic spin columns. A and B PCR purifications were performed according to the respective manufacturer’s recommendations (see Supplementary Material). The PCR purification using proprietary buffers were carried out using generic A’s protocol, with the exception of using 5:1 volume ratio of buffer to PCR reaction (according to generic brand B protocol).

DNA analysis

DNA concentration and A260/280 ratio were analysed spectrophotometrically using IMPLEN Nanophotometer P330 in triplicates. 1% TAE agarose gels were used to analyse quantity and quality of plasmid DNA extracted from the miniprep and PCR purifications. 10 µl of extracted/purified DNA from the above comparisons were loaded with 6x loading dye containing SYBR Green (Quintech Life Sciences) and analyzed using the RunVIEW electrophoresis apparatus (Cleaver Scientific).

Statistical analysis

Time taken for the gel dissolution, DNA concentration and A260/280 ratio from the nucleic acid extractions were analyzed using One-Way ANOVA and independent T-tests. Significance were deemed when p<0.05. All statistical analysis was performed using SPSS 17.0 (IBM).

Results and Discussion

From the systematic testing, we established a set of optimized (OPT) and completely “home-made” (HM) buffers for nucleic acid extraction and purification kits that are comparable to the two generic brands (A and B) in terms of plasmid yield and purity. Through the step-wise buffer substitution (Table 1), we found that P-P1, P-W2-1

P-W1 ----- NH₂C(=NH)NH₂ • HCl, Isopropanol
P-W2-1 ----- Tris Base, Ethanol
P-W2-2 ----- Ethanol
P-EB1 ----- Tris-Base
P-EB2 ----- Tris-HCl, EDTA
P-EB3 ----- NaCl, Tris-Base, Isopropanol

Generic brand A (an “original equipment manufacturer” or “OEM” brand) and generic brand B (well-established brand) buffers were purchased from the commercial vendors.

Culturing of Escherichia coli for miniprep: Luria–Bertani (LB, Biopolis Shared Facilities, BSF, A*STAR) with ampicillin (GoldBio, USA) was used as growth medium. Previously made competent E. coli [4] were transformed with ampicillin resistant plasmids bearing antibody genes as previously described [5], and inoculated in LB ampicillin broth in overnight cultures at 37°C in a shaking incubator. The plasmids were used for miniprep, gel extractions, and PCR amplification. For comparisons, the same plasmids and bacterial cultures were used.

Establishing the OPT and HM buffers with generic brand A miniprep kit: To establish the best “home-made” (HM) proprietary buffers, we evaluated the solution by systematically displacing the buffers in kit A while following its protocol (Supplementary Material). The PCR purification using proprietary buffers were carried out using generic B’s protocol, with the exception of using 5:1 volume ratio of buffer to PCR reaction (according to generic B protocol). Time taken for the gel slices to dissolve completely were measured with a lab timer and analyzed statistically.

Optimization of PCR purification buffers

Polymerase chain reaction: PCR reactions of 325 µl were performed containing 6.5µl of Taq polymerase, 19.5 µl 30 mM MgCl₂, and 32.5 µl of 10X PCR Buffer (Axil Scientific), 26 µl of 2 mM dNTPs mix (Quintech Life Sciences), 13 µl of reverse primer : OriP Nrul R (5’-ATA TCT CGC GAA TGC TGG GGG ACA TGTACC TC-3’), forward primer OriP Nrul F (5’-CAC ACT CGC GAA GGA AAA GGA CAA GCA GCG AA-3’), template plasmid DNA, and 201.5 µl of HyClone water (Thermo Scientific, Cat no. SH30383.01). The amplicon oriP is ~1.9 kb. The completed PCR mix was transferred into PCR tubes of 25 µl aliquots and carried out in Artik Thermal Cycler (Thermo Scientific) with the following profile: Initial denaturation at 94°C for 5 minutes; 30 cycles of denaturation at 94°C for 1 minute, annealing and extension at 71°C for 3 minutes; and final extension at 72°C for 10 minutes.

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DNA concentration and A260/280 ratio were analysed spectrophotometrically using IMPLEN Nanophotometer P330 in triplicates. 1% TAE agarose gels were used to analyse quantity and quality of plasmid DNA extracted from the miniprep and PCR purifications. 10 µl of extracted/purified DNA from the above comparisons were loaded with 6x loading dye containing SYBR Green (Quintech Life Sciences) and analyzed using the RunVIEW electrophoresis apparatus (Cleaver Scientific).

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Table 1: Comparison between the proprietary buffers and generic brand A miniprep kit buffers using DNA yields.

| Buffer type | Buffer comparisons | No of expts | OPT buffer selection | HM buffer selection |
|-------------|--------------------|-------------|----------------------|---------------------|
| Equilibration | No buffer < GA-BK | 3 of 3 | GA-BK | P-BK2 |
| | P-BK1 > GA-BK | 4 of 6 | P-BK1 | P-BK1 |
| | P-BK2 > GA-BK | 2 of 2 | P-BK2 | P-BK2 |
| Resuspension | P-P1 > GA-A1 | 3 of 4 | P-P1 | P-P1 |
| | P-P2 < GA-A2 | 3 of 4 | GA-A2 | P-P2 |
| Lysis | P-P3-1 < GA-A3 | 3 of 4 | GA-A3 | P-P3-2 |
| | P-P3-2 < GA-A3 | 2 of 4 | GA-A3 | P-P3-2 |
| Neutralization | P-W1 = GA-W1 | 2 of 4 | P-W1 | P-W1 |
| | P-W2-1 > GA-W2 | 3 of 4 | P-W2-1 | P-W2-1 |
| | P-W2-2 > GA-W2 | 2 of 4 | P-W2-2 | P-W2-1 |
| Pre-wash | P-EB1 = GA-EB | 2 of 4 | P-EB1 | P-EB1 |
| | P-EB2 > GA-EB | 3 of 4 | P-EB2 | P-EB2 |
| Wash | P-EB3 > GA-EB | 3 of 3 | P-EB3 | P-EB2 |

[DNA] were measured in three separate readings using IMPELN Nanophotometer P330. Independent T-test was used to determine the statistical significance of the differences between test buffers and generic brand A buffers. Differences in [DNA] were measured in three separate readings using IMPLEN Nanophotometer P330. Independent T-test was used to determine the statistical significance of the differences between test buffers and generic brand A buffers. Differences in [DNA] were measured in three separate readings using IMPLEN Nanophotometer P330. Independent T-test was used to determine the statistical significance of the differences between test buffers and generic brand A buffers.
of guanidine would have aided in better adsorption to the silica gels. Being similar to other chaotropic agents (e.g. potassium or sodium iodide), which are necessary for dissolving agarose gels [8], the higher concentrations of guanidine thiocyanate would also dissolve the agarose quicker. Thus, on the basis of timing and yields, P-QG-2 was chosen as the optimal buffer.

For PCR kits, we compared only the PCR binding buffers i.e. the optimized P-W1 and P-QG2 with generic A and B buffers. ANOVA tests showed significant differences between the DNA recovered, $F(5, 48)=261.72, p=0.000$. It was observed that using generic B column, buffer P-W1 (P-W1-B in Figure 3A) obtained the highest DNA recovery, almost up to 30 ng/µL. On the contrary, the same P-W1 buffer in generic A column yielded only slightly above 20 ng/µL, thus supporting previous miniprep observations that generic B spin columns were superior with respect to DNA binding. Normalizing the spin columns by comparing P-W1 on both A and B spin columns, buffers of both generic brands would generate similar yields.

Electrophoresis of the purified PCR products using the different

| Buffer comparisons | Total no of expts |
|--------------------|------------------|
| P-QG2 > Generic A  | 3 of 3           |
| P-QG3 = Generic A  | 3 of 3           |
| P-QG2 > Generic B  | 2 of 3           |
| P-QG3 > Generic B  | 3 of 3           |

Gel Extractions using P-QG2 and P-QG3 were carried out using generic brand A protocol (see Supplementary Material) with the exception of varying the gel dissolving buffer GA-G1. DNA concentrations were measured in three separate readings using IMPLEN Nanophotometer P330. Independent T-tests were used to determine the statistical significance of the differences in DNA recovered. Differences were deemed significant when $p<0.05$

Table 2: Comparison of DNA yields in 3 independent gel extraction experiments using P-QG2 and P-QG3 buffers compared against generic brands A and B buffers using their respective kits and protocols.

Figure 2: Comparison of time taken for excised gel fragments to dissolve completely using generic brand A and B gel dissolving buffers, P-QG2 and P-QG3 buffer.

Bar chart showing the means and standard errors of the time taken for excised gel fragments to dissolve completely using OPT buffers of generic A and B. P-QG2-A and P-QG2-B buffer testing were carried out using brand A’s protocol (see Supplementary Material). On the other hand, P-QG2-B and P-W1-B buffers were performed on brand B’s spin column (without P-BK).

Figure 3A: Comparison of the DNA recovered using P-QG2, P-W1, generic brands A and B’s PCR purification buffers.

Bar chart showing the means and standard errors of the DNA recovered using various buffers. P-QG2-A and P-W1-A buffer testing were carried out using brand A’s protocol (see Supplementary Material). On the other hand, P-QG2-B and P-W1-B buffers were performed on brand B’s spin column (without P-BK). DNA concentrations were determined by IMPLEN Nanophotometer P330 in three separate extractions measured in triplicates.

Analysis of the P-QG2 and P-W1 recipes found that higher concentrations of guanidine (by ~1 M in P-QG2) resulted in better purification, which we propose to result from the release of polymerases from DNA, allowing their adsorption to the silica.

As a final comparison, we carried out trials comparing the full set of HM buffers and OPT buffers against both generic A and B (Figure 1). As can be observed, OPT buffers had the best yields regardless of the columns used, with HM buffers comparable to the commercial brands A and B.

Conclusion

The findings of the study allowed us to rely on more cost-effective columns without compromising experiments. Extending beyond the kits tested, the factors of these buffers also underline processes such as midi, maxi and giga scale DNA extractions, allowing labs to optimize their own cost-effective reagents by the addition of important chemicals to their existing buffers or kits (e.g. adding a Na+ column equilibration step to existing commercial kits). Through detailed analysis of buffer constituents, we were able to validate the importance of:

1) Na+ concentrations in column equilibration.

2) Importance of strong acids for low pH in the neutralization of cell lysis buffer.
3) High salt for higher stringency in column washes.
4) Tris and chelating agents to remove nuclease cofactors and pH buffers that would not generate free radicals.
5) High presence of chaotropic agents for faster gel dissolution and removal of interfering proteins for both gel and PCR kits.

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Competing Interests
This work was commissioned by Quintech Life Sciences Pte Ltd to explore the factors to making better buffers, which may be made commercially available. There are no other competing interests.

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Figure 3B and 3C: Comparison of the P-QG2 and P-W1 with generic brand A and B PCR purification kits.
(B) Comparisons of P-QG2 and P-W1 buffers were performed using generic A spin columns.
(C) Comparisons of P-QG2 and P-W1 buffers were performed using generic B spin columns.
10uL of purified PCR DNA were mixed with 6x loading dye and loaded on a 1% TAE agarose gel using RunVIEW by Cleaver Scientific. Samples were processed from the same PCR reaction for consistency.