Decreasing formalin concentration improves quality of DNA extracted from formalin-fixed paraffin-embedded tissue specimens without compromising tissue morphology or immunohistochemical staining

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ABSTRACT

Genomic technologies are increasingly used clinically for both diagnosis and guiding cancer therapy. However, formalin fixation can compromise DNA quality. This study aimed to optimise tissue fixation using normal colon, liver and uterus (n=8 each) by varying neutral buffered formalin (NBF) concentration (1%–5% w/v) and fixation time (24–48 hours). Fixation using 4% NBF improved DNA quality (assessed by qPCR) compared with routine (4% unbuffered formal saline-fixed) specimens (p<0.01). Further improvements were achieved by reducing NBF concentration (p<0.00001), whereas fixation time had no effect (p=0.110). No adverse effects were detected by histopathological or QuPath morphometric analysis. Immunohistochemistry for multicytokeratin and α-smooth muscle actin revealed no changes in staining specificity or intensity in any tissue other than on liver multicytokeratin staining intensity, where the effect of fixation time was more significant (p=0.0004) than NBF concentration (p=0.048). Thus, reducing NBF concentration can maximise DNA quality without compromising tissue morphology or standard histopathological analyses.

INTRODUCTION

The increasing use, accessibility and cost-effectiveness of genomic analyses for both clinical and research purposes promise to revolutionise cancer medicine. However, the routine histopathological tissue from which DNA is extracted is typically formalin-fixed and paraffin-embedded (FFPE) and notoriously variable in the quality of the nucleic acids extracted.1 DNA extracted from FFPE tissue is fragmented, crosslinked and contains abasic sites. This can impact on PCR-based sequencing assays, where the stochastic effects of low template copy number can lead to false mutation calls, particularly when combined with a low tumour cell content/low percentage mutation.2 Furthermore, there are the additional concerns of toxicity and carcinogenicity associated with formalin use.3 Endeavours have focused on finding alternative, ‘molecular’ fixatives but the widespread use of these agents has broadly failed to gain any traction over standard formalin-based methods that are used globally in clinical laboratories. This is partly due to expense and the fact that re-optimisation of immunohistochemistry (IHC) protocols is usually required and, in general, formalin outperforms other fixatives for most antibodies tested.4

Although it is well established that longer formalin fixation times reduce DNA yield/quality, the effects of reducing formalin concentration on DNA quality or tissue morphology have received scant attention.5 This study aimed to optimise formalin fixation protocols with a view to maximising extracted DNA quality by varying both formalin concentration and fixation time, as well as investigating the effects on tissue morphology and IHC staining.

MATERIALS AND METHODS

Specimen collection and fixation

Anonymised specimens were collected immediately after surgery and included colon, liver and uterus (n=8 for each). All materials collected were normal background tissues from resections surplus to diagnostic requirements. Tumour tissue was deliberately excluded in order to obviate the impact of variable tumour necrosis across malignant specimens. Neutral buffered formalin (NBF) solutions were made by diluting 40% w/v formaldehyde (Solmedia, Shrewsbury, UK) in phosphate buffer solution (PBS). Tissue samples were divided into 10 pieces (<5 mm thickness) and fixed under the following conditions: 5%, 4%, 3%, 2% and 1% (w/v) formaldehyde in PBS for 24 and 48 hours at room temperature, after which they were placed into 70% ethanol prior to routine embedding in paraffin.5 Given the estimated penetration constant of formalin,6 complete fixation would have been achieved at both time points. In parallel, tissue-matched specimens (n=4 each) fixed according to our routine clinical service protocol (10% unbuffered formal saline; 4% w/v formaldehyde at room temperature for 24–48 hours) were collected in order to assess the adequacy of our routine tissue fixation regimens.

DNA extraction and assessment of DNA quality

DNA was extracted from tissue sections using a QIAamp DNA FFPE Tissue Kit (Qiagen, Manchester, UK) and included an RNase step. The DNA concentration was determined on a
Figure 1  Effect of formalin fixation conditions on DNA quality. (A) Comparison of quality of DNA extracted from experimentally fixed FFPE tissues (n=8 each, fixed using neutral buffered formalin (4% w/v formaldehyde) for 24 hours; closed circles) with that extracted from routinely fixed specimens (n=4 each, fixed using unbuffered formal saline (4% w/v formaldehyde); open circles). Graph depicts amplifiable copy number (median and IQR) of a 180 bp FTH1 fragment, measured by qPCR, both in absolute terms (right y axis) and relative to fresh human genomic DNA (extracted from whole blood), seri-

NanoDrop ND-1000 spectrophotometer (ThermoFisher Scientific, Loughborough, UK). Quantitative PCR using primers amplifying an 180bp region of the FTH1 gene (TaqMan assay ID hs01694011-m1) was used to assess amplifiable copy number. Duplicate reactions were performed using 50ng input DNA in TaqMan Gene Expression Master Mix on a QuantStudio 5 Real-Time PCR System (ThermoFisher Scientific). Amplifiable copy number was determined using a standard curve prepared from the effective (PCR-amplifiable) DNA copy number, potentially leading to stochastic effects on next-generation sequencing (NGS) results. We therefore chose to assess DNA quality using a qPCR-based approach.9 We began by comparing DNA quality from our experimentally fixed colon, liver and uterine tissue samples (fixed for 24 hours in standard 10% NBF; 4% w/v formaldehyde) with contemporaneous samples fixed in the routine histopathology laboratory using unbuffered formal saline (4% w/v formaldehyde) (figure 1A). There were no significant differences in DNA quality between the different tissue types in either fixation condition. However, for each tissue type, the quality of DNA was significantly higher in experimentally fixed compared with routinely fixed samples, where none of the former and half of the latter had fewer than 100 amplifiable copies per 50ng DNA input (equivalent to <0.6% of the amplifiable copy number of fresh genomic DNA). We next examined the effect of varying NBF concentration on the quality of extracted DNA (figure 1B). Analysis of pooled copy number data from all tissues (n=24 for each fixation condition) showed the overall effect of NBF concentration on DNA quality was highly significant (p=1.63×10^-9), whereas the effect of fixation time was not (p=0.110). There was no significant interaction between the two variables (p=0.609); the Scheirer-Ray-Hare test. Post hoc testing revealed significant increases in amplifiable

H&E staining and IHC  

Tissue sections (5 μm) were mounted on Superfrost Plus slides and H&E stained using standard protocols. IHC was carried out with antibodies against multicytokeratin (MCK; AE1/AE3; PA909) and α-smooth muscle actin (SMA; αSM-1; PA0943), both Leica Biosystems, Milton Keynes, UK, using the Leica Bond III autostainer IHC validated protocol F, with the Bond Polymer Refined Detection (DS9800). Slides were digitised at 20× magnification on Aperio XT slide scanners (Aperio Technologies, Vista, California, USA) with a compression rate of 70% and the images hosted on the University of Leeds servers. A review of H&E digital images was performed blind by independent scorers (NMO, KA) and assessed against predetermined criteria including nuclear morphology, architectural integrity, presence of retraction artefact and appropriateness of H&E staining. Slides were scored 2, 1 and 0 indicating good, sub-par and compromised morphology for diagnostic purposes, respectively.1 Morphometric assessment of H&E images was carried out using QuPath,8 using the Watershed Nuclear Detection algorithm to measure both numbers of nuclei per unit area and mean nuclear area, selecting a minimum of 1 million µm^2 per slide for analysis (glandular and myometrial areas were selected for colon and uterus, respectively). In order to obviate interobserver variability in IHC scoring, staining intensity and specificity were also measured objectively using QuPath. Colour deconvolution was applied followed by automated tissue detection. Simple Linear Iterative Clustering superpixels were then calculated for the annotations for each image. 3,3'-Diaminobenzidine (DAB) optical density (OD) intensity features (mean, minimum, maximum and median) were then added to the superpixels at a width of 25 μm. A thresholding script was then used to delineate positive (stained) from negative (unstained) superpixels based on the mean DAB OD for each antibody–tissue combination. Median DAB OD value for each categorised superpixel was exported and the median values of all positive and negative superpixels per slide calculated.

Statistics  

Statistical analysis was performed using R or IBM SPSS. Comparisons were made by non-parametric tests (Mann-Whitney U, Kruskal-Wallis, Scheirer-Ray-Hare and χ² tests), as appropriate. Dunn post hoc tests with Benjamini-Hochberg false discovery rate (FDR) correction for multiple comparisons were also performed, as appropriate.

RESULTS  

Effect of formalin fixation conditions on DNA quality  

Formalin fixation causes a number of artefacts that decrease the effective (PCR-amplifiable) DNA copy number, potentially leading to stochastic effects on next-generation sequencing (NGS) results. We therefore chose to assess DNA quality using a qPCR-based approach.9 We began by comparing DNA quality from our experimentally fixed colon, liver and uterine tissue samples (fixed for 24 hours in standard 10% NBF; 4% w/v formaldehyde) with contemporaneous samples fixed in the routine histopathology laboratory using unbuffered formal saline (4% w/v formaldehyde) (figure 1A). There were no significant differences in DNA quality between the different tissue types in either fixation condition. However, for each tissue type, the quality of DNA was significantly higher in experimentally fixed compared with routinely fixed samples, where none of the former and half of the latter had fewer than 100 amplifiable copies per 50ng DNA input (equivalent to <0.6% of the amplifiable copy number of fresh genomic DNA). We next examined the effect of varying NBF concentration on the quality of extracted DNA (figure 1B). Analysis of pooled copy number data from all tissues (n=24 for each fixation condition) showed the overall effect of NBF concentration on DNA quality was highly significant (p=1.63×10^-9), whereas the effect of fixation time was not (p=0.110). There was no significant interaction between the two variables (p=0.609); the Scheirer-Ray-Hare test. Post hoc testing revealed significant increases in amplifiable

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copy number in samples fixed with 1% and 2% formaldehyde compared with those fixed with the standard NBF solution (4% formaldehyde) at both time points. Additionally, there was a significant difference between samples fixed in 3% vs 4% formaldehyde at 48 hours (figure 1B). Although a third of samples fixed for 48 hours in the standard NBF solution had <10% of the amplifiable copy number of fresh DNA, all of the samples fixed with 1% formaldehyde for the same length of time were above this threshold.

Effect of formalin fixation conditions on tissue morphology and IHC staining

When H&E sections were blind reviewed by two histopathologists, no significant differences in tissue morphology scores were identified between different fixation conditions in any tissue (figure 2 and table 1).

Conclusions

The experimental FFPE samples fixed for 24 hours with the standard concentration (4% w/v) of NBF yielded significantly better quality DNA than those fixed in the routine diagnostic histopathology laboratory in the standard concentration of unbuffered formal saline (a practice no longer used in our clinical laboratory). This agrees with the majority of studies using a PCR-based

Table 1: Histopathological assessment of the effects of buffered formalin fixation conditions on colon, liver and uterine tissues

| Tissue | Colon (p=0.628) | Liver (p=0.609) | Uterus (p=0.427) |
|--------|----------------|----------------|------------------|
| Fixation | 1 (n) 2 (n) 1 (n) 2 (n) 1 (n) 2 (n) | 1% to 24 hours | 2% to 24 hours | 3% to 24 hours | 4% to 24 hours | 5% to 24 hours |
| 1% to 24 hours | 3 | 5 | 1 | 7 | 0 | 8 |
| 2% to 24 hours | 4 | 4 | 1 | 7 | 0 | 8 |
| 3% to 24 hours | 5 | 3 | 0 | 8 | 1 | 7 |
| 4% to 24 hours | 5 | 3 | 0 | 8 | 0 | 8 |
| 5% to 24 hours | 3 | 5 | 0 | 8 | 0 | 8 |
| 1% to 48 hours | 5 | 3 | 1 | 7 | 0 | 8 |
| 2% to 48 hours | 4 | 4 | 0 | 8 | 0 | 8 |
| 3% to 48 hours | 1 | 7 | 0 | 8 | 0 | 8 |
| 4% to 48 hours | 3 | 5 | 0 | 8 | 0 | 8 |
| 5% to 48 hours | 4 | 4 | 0 | 8 | 0 | 8 |

H&E-stained sections (n=8 cases for each tissue) were blind reviewed by two histopathologists and given scores (0–2; no scores of zero were given). P-values for each tissue type are indicated (χ² test).
assay to estimate DNA quality. A likely cause is the oxidation of formaldehyde to formic acid, which in unbuffered fixative leads to reduced pH, resulting in DNA depurination and consequently reduced PCR amplification efficiency.

This study showed that reducing the formaldehyde concentration in buffered formalin fixative further improved DNA quality, whereas varying time had no significant effect on the time points used in this study (24 and 48 hours). The majority of PCR-based studies agree that fixation times of <72 hours are preferable for maximising DNA integrity. Although the improvements afforded by reducing formaldehyde concentration are less dramatic than those observed when switching from unbuffered to buffered formalin fixative, they are still meaningful from a clinical perspective. For example, a third of samples fixed for 48 hours in standard (4%) buffered formaldehyde have a copy number of <10% that of fresh DNA, equivalent to 333 copies per 10 ng input, whereas all of the samples fixed for the same time in 1% and 2% formaldehyde exceeded this threshold. Reported threshold copy number input for accurate NGS-based mutation quantification or detection were 379 and 95, respectively, for mutation frequencies of 30%–40%.

Reducing formaldehyde concentration had no appreciable effect on histological architecture, nuclear morphology or quality of H&E staining. Morphometric analysis using QuPath revealed no significant effects on either the mean nuclear area or the number of nuclei per unit area (as a measure of tissue shrinkage). Surprisingly, very few studies addressing the effect of formaldehyde concentration on tissue morphology/morphometry have been published. Nonetheless, these have shown has the potential to increase the number of samples amenable to accurate mutation testing, especially those with low cellular content/percentage tumour cells/mutation frequency.
that reducing formaldehyde concentration to 1%–2% had no substantial effect on morphology\textsuperscript{14} or morphometry,\textsuperscript{15} although these studies lacked statistical analysis.

Changing the NBF fixation conditions had no visible effect on IHC staining intensity or specificity with MCK and SMA antibodies when reviewed blind by two histopathologists. Although QuPath analysis detected a significant effect of formaldehyde concentration on liver MCK staining intensity, the effect of fixation time was more significant overall. Moreover, there were no measurable differences in staining specificity in any antibody-tissue combination tested. These observations are in line with the fact that the basic chemistry of fixation is unchanged unlike with alternative (non-crosslinking) fixatives, where antibodies require systematic re-optimisation of antigen-retrieval protocols and in some instances, only work on FFPE specimens.\textsuperscript{3,16} Thus, reducing the formaldehyde concentration in buffered formalin fixatives has the potential to increase DNA quality and reduce occupational exposure to formaldehyde while avoiding these substantial barriers to clinical implementation.

**Take home messages**

- Reducing the formaldehyde concentration in buffered formalin fixative to 1%–2% significantly increases DNA quality without compromising tissue morphology or immunohistochemical staining.
- Implementation by diagnostic histopathology laboratories would be relatively straightforward and would increase the number of samples amenable to genomic analysis while also reducing occupational exposure to formaldehyde.

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