Mammographic compression practices of force- and pressure-standardisation protocol: A scoping review

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Keywords
breast, mammographer, mammography, review, screening

Abstract

Introduction: As an efficient, effective and moderately inexpensive modality, mammography has been implemented as a cancer screening tool and in diagnostic management. However, appropriate breast compression is necessary for optimal outcomes. Current key measures of compression force are subjective and variable, giving rise to the concept of a 'personalised' pressure-standardisation protocol. Methods: A scoping review of the literature was performed using the Arksey and O’Malley framework to explore the existing force- and pressure-standardisation protocols in clinical application. A comprehensive search strategy and standardised study selection and evaluation were completed. This synthesis of existing knowledge can lead to the implementation of mechanically standardised mammographic compression pressure as a feasible tailored approach to clinical practice. Four databases (PubMed, MEDLINE, Embase and Scopus) were searched from the databases’ inception to 13 December 2019 for relevant information, and eighteen articles were selected for analysis. Results: In addition to current protocol comparison, emerging key concepts include the reasoning behind standardisation, the benefits of improved diagnostic outcomes/decreased pain with negligible change in image quality and average glandular dose (AGD), and the recommendation of a 10kPa (approximate) pressure-standardisation protocol. Research to date is largely based abroad (Netherlands), with a strong focus on screening practices. Consequently, several gaps in the current literature were identified as potential directions for future investigation. Conclusions: As a suggested mammographic guideline, compression pressures of approximately 10kPa aid in image acquisition reproducibility both within and between women; pain levels decrease, with minimal variations to breast thickness, AGD and image quality.

Introduction

As breast cancer is one of the most prevalent female cancers, it is unsurprising that mammographic screening programs have been implemented worldwide.1 Such programs are estimated to result in an approximate annual reduction in breast cancer mortality by 30%.2 Additionally, mammography is a cost-effective and gold standard practice.3 Sources report high sensitivity and the highest specificity in mammographic screening relative to MRI; higher MRI sensitivity does not translate into improved clinical outcomes due to the ineffective detection of ductal carcinoma in situ.3 Furthermore, mammography is the preliminary imaging modality used in most diagnostic settings (depending on patient age); calcium deposits, most frequently benign, can be indicative of potentially cancerous pathological change, which are usually seen clearly on a mammogram.4,5 Breast compression is necessary for any successful mammographic examination; many sources discuss the effect of a uniform breast tissue thickness on improving image contrast and quality, with an associated reduction
in radiation dose, geometric/motion blurring and tissue superimposition.\textsuperscript{1,3–14} Whilst ultimately aiding in the detection of pathology, the discomfort attributed to breast compression remains the highest reason for patient non-compliance with screening programs and lack of diagnostic lesion visibility.\textsuperscript{9,15,16} According to the literature, the key factors affecting compression can be attributed to the patient, the mammographer and/or the equipment.\textsuperscript{17} However, compression standardisation is difficult due to the lack of explicit criteria to assess consistency; the predictability and reproducibility of mammographic examinations are thus difficult to determine.\textsuperscript{13} This is increasingly apparent when comparing the mammography protocols utilised on a global scale, in conjunction with international approaches to clinical practice – the specification of concrete parameters and/or guidelines is brief and minimal, with sufficient compression seemingly based on mammographer opinion and experience. The only objective measures currently obtainable in real time are those of compression force and breast thickness; these are gained mechanically from the mammography system, yet do not account for individual variations in breast size and elasticity.\textsuperscript{13} This has led to recent investigation of pressure (expressed in kilopascals – kPa) as a ‘personalised’ adaptation of force. Since pressure considers both the force and contact area simultaneously (pressure = compression force/breast contact area), it has been reported as a more physiologically appropriate compression parameter than force.\textsuperscript{14} In comparison, an opinion-based application of force is variable and unreliable, whilst a mechanical application of target pressure is consistent and reproducible. Consequently, recent research has stemmed into developing an objective pressure-standardisation protocol for clinical implementation, which allows for real-time appraisal. Therefore, the aim of this review is to assess the feasibility of mechanically standardised mammographic compression in clinical practice. This will be evaluated in comparison with current techniques – both nationally and internationally – with the perceived benefits, recommendations and limitations outlined.

**Methods**

**Method of review**

A scoping, or ‘mapping’, review was performed to assess the current standing of mammographic compression practices available in existing literature. The organisational framework, derived as described by Arksey and O’Malley, was chosen as a means of evaluating the extent of available evidence for a clinical overview.\textsuperscript{18} As information from existing evidence was extracted and summarised for this scoping review, rather than methodologically appraised and statistically analysed, a systematic review was deemed unsuitable. To establish the context of this research in the wider mammographic community, the following research questions were pondered:

- Would mammographic practices benefit from mechanical standardisation of objective parameters?
- In what ways can compression force and compression pressure be justified as an appropriate measure of standardisation?
- How are outcomes relevant from both a clinical and patient-centred perspective?

**Search strategy**

Four electronic databases (PubMed, MEDLINE, Embase and Scopus) were searched for English-language articles discussing mammographic compression practices from the databases’ inception to 13 December 2019; most sources were found to be relatively recent due to the nature of the topic. Structured search strategies were devised to optimise findings; the initial search began with the Medical Subject Heading (MeSH) term ‘mammography’, which was altered to ‘mammograph*’ as a database entry. Further keywords added to the search strategy included (AND) ‘compress*’ OR ‘force’ OR ‘pressure’, AND ‘standard*’. Although no relevant MeSH terms exist for such keywords, these were deemed relevant to the aims of the research. Search fields were set at ‘all fields’ for each database, except for Scopus, which was limited to ‘article title, abstract, keywords’. The number of hits from each database was 53, 35, 43 and 60 articles from Scopus, PubMed, MEDLINE and Embase, respectively.

Following the addition of hand-sought records from expert contribution and duplicate removal by a single researcher, titles and abstracts were screened independently by two researchers. The independent screening and reviewing of eligible studies were in line with the 2005 scoping review framework by Arksey and O’Malley,\textsuperscript{18} as well as the Preferred Reporting Items for Systematic Review and Meta-Analysis (PRISMA)\textsuperscript{19} guidelines. Specified eligibility criteria for each step of the selection process were adhered to systematically, with eligible articles then selected by reading the full text. Eligible titles were those relevant to ‘mammography’ or ‘breast’, and eligible abstracts discussed at least one aspect of ‘compression’, ‘force’ or ‘pressure’. Review articles, case reports, conference reports, letters, editorial comments, opinions (including qualitative reports) and non-English articles were excluded. Other exclusion criteria included articles purely discussing quality assurance/physics without reference to clinical...
applicability. Any disagreement was discussed and resolved by consensus among the entire research team. The team comprised of researchers with a medical imaging background, including an experienced breast imaging specialist with post-graduate qualification in breast imaging. The entire research team also had extensive experience in conducting scoping reviews, systematic reviews and meta-analyses.

**Literature analysis**

A data extraction form was developed using Microsoft Excel to collect information related to (i) country of origin and current practices, (ii) the perceived need for standardisation, (iii) benefits, (iv) recommendations and (v) limitations. Information compiled in the extraction form was reviewed by two researchers, with data analysed for key concepts and frequencies. These included overall benefits of the standardisation protocols, such as diagnostic outcomes, image quality, AGD and patient experience. A total of 18 articles were included for analysis in this scoping review; the PRISMA flow chart (Figure 1) details the review process, whilst Table 1 presents the summary of the reviewed studies, arranged alphabetically:

**Results**

The retained articles consisted of a mixture of quantitative study methods. There were six prospective and eleven retrospective studies, with one
Table 1. This table presents the data extraction and analysis of the selected studies.

| Authors                       | Country of origin | Protocol used | Study details                                                                 | Why standardise?                                                                                   |
|-------------------------------|-------------------|---------------|--------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------|
| Branderhorst et al.            | Netherlands       | Netherlands: Dutch screening QA protocol – 12daN. USA: no target – taut breast, unless painful | Retrospective cohort: Netherlands: screening asymptomatic patients, aged 50–75; CC + MLO USA: screening/diagnostic (symptomatic/asymptomatic), aged 50–75; CC + MLO | Account for contact area; contact area decreases, average pressure/variance increase, despite lower force |
| Branderhorst et al.            | Netherlands       | Prospective cohort: Real-time and retrospective methods/software studied on asymptomatic patients; CC + MLO | | Ensure contact area between breast/paddle can be determined for accurate pressure readings          |
| de Groot et al.                | Netherlands       | Compress to 18daN or until pain threshold | Prospective observational: Hospital setting excluding interventions, patients aged 30–88; CC + MLO | Force-controlled compression causes large pressure variation; pressure-controlled accounts for contact area as predictive parameter for severe pain |
| de Groot et al.                | Netherlands       | 18daN compression protocol | Prospective observational: Real-time examination of post-intervention (unilateral) patients | Protocol applicable to post-conservation therapy population |
| de Groot et al.                | Netherlands       | Netherlands: Dutch screening QA protocol – 12daN. USA: no target – taut breast, unless painful | Retrospective cohort: Netherlands: screening asymptomatic patients, aged 50–75; CC + MLO USA: screening/diagnostic (symptomatic/asymptomatic), aged 50–75; CC + MLO | Account for contact area; contact area decreases, average pressure/variance increase, despite lower force |
| de Groot et al.                | Netherlands       | 14daN force standardisation, to be compared with 10kPa pressure standardisation | Prospective cohort: Screening asymptomatic patients aged 50–75; CC + MLO | Force applied in proportion to contact area (account for breast size) |
| de Groot et al.                | Netherlands       | 18daN force standardisation, compared to 10kPa standard pressure | Case-control: Case pairing of patients over time, with alternate protocol used for each; CC + MLO | Ensure lesion appearance is consistently satisfactory |
| den Boer et al.                | Netherlands       | Compress to 100–150N or until tolerance | Retrospective cohort: Screening follow-up of asymptomatic patients, mean age 59; CC + MLO (separate) | Compression as a function of contact area, to apply pressure independent of breast size |
| Holland et al.                 | Netherlands       | Retrospective cohort: Screening asymptomatic patients, aged 50–75; MLO views only | Possible improvement to screening program performance |
| Holland et al.                 | Netherlands       | Retrospective cohort: Screening asymptomatic patients, aged 50–75; MLO only | Possible improvement to screening program performance |
| Jeukens et al.                 | Netherlands       | 10kPa standard compression | Retrospective cohort: Screening follow-up of asymptomatic patients, mean age 59; CC view | Reduce pain due to compression |
| Lau et al.                     | Malaysia          | Compress until taut or intolerable pain (no target force) | Retrospective cohort: Screening/diagnostic imaging of symptomatic/asymptomatic patients, aged 35–80 (clinical study); CC + MLO | Compression based on contact area; optimise protocols for Asian women (assumed to have smaller breasts) |
| Mercer et al.                  | United Kingdom    | Subjective mammographer judgement | Retrospective cohort: Evaluating mammographer practice in screening service | Minimise compression practice variance between/within mammographers |
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## Benefits of Pressure Standardisation

| Diagnostic outcome | Image quality | AGD | Patient experience | Recommendations | Limitations |
|--------------------|---------------|-----|-------------------|----------------|-------------|
| Reduced pain       | 10kPa standard pressure; reduced variation and clear standards | Screening vs diagnostic cohort |
| Reduced pain       | 10–12kPa standard pressure; increased thickness of 9% (small) and 2% (large) breasts | Clinical only – not representative of screening population |
| Reduced pain post-intervention | Support for 10kPa standard pressure | Post-intervention only |
| Potentially reduced pain | 10kPa standard pressure | Screening vs diagnostic cohort |
| Not compromised | Similar between protocols | Screening only – not extend conclusions to lesion detection |
| Unchanged | Reduced pain: 1/2 cohort = less pain, 1/3 = unchanged | Changes in practice between examinations |
| Unchanged | Reduced unnecessary pain | |
| Not compromised, more consistent | Reduced pain | Time delay between successive acquisitions |
| Moderate pressures best (statistically insignificant) | Unchanged | Reduced pain | Quantitative protocol; high compression worse on lesion visibility than low | No CC view |
| High pressure decreases sensitivity; low pressure decreases specificity | Unchanged | Reduced pain | No optimal pressure; high compression worse on lesion visibility than low | No CC view; international screening differences; pectoral interference |
| No clinically relevant difference | No clinically relevant difference | None of clinical relevance | Psychological impact on post-intervention/follow-up examinations |
| Not compromised | Potentially reduced pain | Compression reduction of 32.5% from 12–9daN | Assume Asian women have small breasts, and study phantom = human tissue |
| Improved consistency | Improved consistency | Varied force impact screening attendance | Focus on training process |
| Improved consistency | Consistently applied force per patient, regardless of mammographer | Small, single-centre study |
Table 1. (Continued)

| Authors          | Country of origin | Protocol used                        | Study details                                                                 | Why standardise?                              |
|------------------|-------------------|---------------------------------------|-------------------------------------------------------------------------------|-----------------------------------------------|
| Mercer et al.16  | United Kingdom    | Subjective mammographer judgement     | Retrospective cohort; Evaluating mammographer practice in screening service    | Minimise compression practice variance between/within mammographers |
| Moshina et al.1  | Norway            |                                       | Retrospective cohort; Screening asymptomatic patients aged 50–69; CC + MLO   | Optimise the performance of screening measures |
| Moshina et al.22 | Norway            | 10kPa standard compression            | Prospective cohort; Screening asymptomatic patients aged 50–69; CC + MLO     | Minimise pain experienced during screening mammography |
| Poulos & McLean9 | Australia         | ‘breast is taut at the sides’; ‘skin blanches’ | Prospective cohort; Based on participants in screening setting                 | More objective criteria for application of compression force       |

Discussion of findings

This study aimed to provide a structured insight into the potential benefits of implementing mechanically standardised mammographic compression pressure in clinical practice. Findings are assessed in the following discussion alongside current practices and directions for suggested future research.

Current practices

The subjectivity and lacking consistency of current compression guidelines are key concerns raised in much of the current literature; many studies describe the variation in applied compression force among mammographers, screening centres and countries.1,5,6,10,13–16,20,24 Clinically, force is typically adjusted in an experience-based manner according to breast size, elasticity and pain threshold; 8,10,13,20,21 Mercer et al. suggest that variation in compression may be associated with technique adaptation, as opposed to inconsistent practice.17 These parameters will differ significantly in a given population; thus, whilst conventional methods attempt to minimise disparity with the implementation of a target force, this is independent of individual breast characteristics.6,8,13 Further to this, it is noted that the compression forces used during mammographic examinations are attributed to the mammographer, as opposed to the patient.15,16 Current European guidelines propose that ‘the breast should be properly compressed, but no more than is necessary to achieve a good image quality’;11 this corresponds with a maximum compressive force of 200N.10 Likewise, the United States of America (USA) recommendations suggest that the breast be compressed until ‘taut’ or ‘just less than painful’,10,13 whilst studies conducted in Malaysia utilise similar subjective techniques.24 Dutch screening programs operate with a force compression protocol, with targets ranging between 12–18 daN.5,8,10,13,14,21 A study completed by Branderhorst et al. comparing the mammographic compression practices within the USA and Netherlands found targets (mean ± standard deviation) to be 13.7 ± 5.9kPa versus 8.1 ± 4.1kPa, and 13.8 ± 2.7 daN versus 7.4 ± 3.1 daN for the Netherlands and USA, respectively.11 Overall, large variations in force, and even larger variations in pressure, were listed within
and between both data sets, with distinctly higher compression limits in the Dutch data.\textsuperscript{10,13} The variations in applied force and pressure during routine mammographic compression are supported with a phantom study completed by Lau et al.\textsuperscript{24} Regarding Australian practices, no formalised documentation of national protocols could be found in the literature, aside from stating the maximum operable force on a mammographic unit; this is 200N for motorised force and 300N for manual force.\textsuperscript{25} However, the subjective criteria of breast tautness and skin blanching are widely accepted.\textsuperscript{9} Overall, this seems to highlight a clinical need for standardised protocol.

The need for standardisation
Pressure standardisation operates on the premise of compression force applied by the paddle to the breast, divided by the contact area between the paddle and the breast.\textsuperscript{6–8,13,14} By definition, it is most probable that pain experienced upon compression is more closely correlated with pressure than force.\textsuperscript{6} For completeness, Branderhorst et al. confirm that the contact area between the breast and compression paddle can be accurately and precisely determined with existing technology.\textsuperscript{20} Current literature suggests pressure is a ‘personalised’ version of force,\textsuperscript{14} with relevance to physiological factors such as tissue composition and blood pressure.\textsuperscript{6,14} The theoretical dependence on contact area is of particular note; the same force results in a higher pressure when applied to a small contact area (i.e. small breast) than a large contact area (i.e. large breast).\textsuperscript{1,6–8,10,13,14,20,21,24} The Dutch study conducted by Branderhorst et al. described a trend between force and contact area; average pressure and variance increases as contact area decreases.\textsuperscript{13} This was found to correlate with pain severity; patients with small breasts experience severe pain more often than large breasts when force standardisation is used.\textsuperscript{14} A Dutch study conducted by de Groot et al. reached similar conclusions; an intra-individual comparison of 14daN force standardisation and 10kPa pressure standardisation was found to decrease pain severity without compromising image quality.\textsuperscript{14} A Norwegian study conducted by Moshina et al. extended these findings, concluding that pressure standardisation could optimise the performance of early screening measures.\textsuperscript{1} Ultimately, pressure-standardisation protocols satisfy the need for objective criteria in mammographic breast compression; this would offer concrete mammographer guidelines, whilst consistently adjusting for individual breast parameters, such as size and elasticity.

Benefits of pressure standardisation
The diagnostic outcomes of pressure-standardised breast compression have been statistically evaluated in recent research. Generally, it was found that excessive pressure decreases mammographic sensitivity, whilst insufficient pressure decreases mammographic specificity.\textsuperscript{6} This reflects the findings described by Moshina et al., whereby a high compression force (directly proportional to pressure) decreased both sensitivity and specificity.\textsuperscript{1} The apparent conflict between the specificity outcome can be attributed to factors influencing resultant image contrast

| Benefits of pressure standardisation | Recommendations | Limitations |
|------------------------------------|----------------|-------------|
| Diagnostic outcome | Image quality | AGD | Patient experience | Establish guidelines for cessation of compression force | Geographical proximity implies similar training |
| Potential improvement in cancer detection | Potential positive impact | Potential reduction | Stabilisation may increase re-attendance | Force of at least 130N, pressure less than 9.8kPa | Patient positioning of questionable quality |
| High force (proportional to pressure) decreases sensitivity and specificity | Possible improvement (inconclusive) | Increased with pressure-standardised paddle (inconclusive) | Higher pain with fixed paddle – clinically inconclusive | None of clinical relevance | Subjectivity of data acquisition |
| Reduced with pressure-standardised paddle (inconclusive) | | | | | |

Compliance with Ethical Standards

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at the extreme ends of the compression scale. Also, Moshina et al. basis conclusions on force alone, independent of contact area.

This is important to note as results cannot be aligned with measures of compression pressure.

Current literature concludes that image quality remains unchanged with a pressure-standardisation protocol; no studies have shown otherwise. De Groot et al. investigated this trend explicitly, noting that although breast compression is milder with a pressure-standardisation protocol, the visibility, contrast and sharpness of stable lesions remain virtually unchanged. Another study even proposed that a force reduction of as much as one third is possible with minimal impact on image quality. As noted by recent research, a large range of pressures result in diagnostically acceptable images in the digital setting; a minimum must exist, though, as image quality is obviously degraded without compression. Nevertheless, Holland et al. suggest over-compression may diminish image quality to a greater extent than under-compression; lesion detectability is compromised as suspicious densities are dissolved. However, this primarily correlates with the high pressures required for spot compression. The opposite can likewise be reasoned; too little compression degrades image quality in that the potential for small cancers to remain undetected increases.

The amount of radiation dose received by the patient with a shift in compression protocol was also investigated. Overall, it could be concluded that pressure standardisation has a negligible impact on AGD, despite the slight increase in breast thickness. Whilst the literature does demonstrate variation in data surrounding the actual dose received, this may possibly be attributed to different system types and automatic exposure control (AEC) settings. The proportion of image repeats was also found to remain unchanged.

Although another recurring issue associated with breast compression is the patient experience of pain, a pressure-based protocol was found to reduce unnecessary pain. This has the potential to positively impact compliance with screening programs and benefit the post-intervention patients for whom follow-up mammograms are mandatory. De Groot et al. revealed that the implementation of pressure standardisation resulted in half the cohort reporting less pain and a third reporting no difference when compared to force standardisation. However, this appears to apply to protocols in entirety; studies explicitly comparing force- and pressure-controlled compression paddles found no preferential basis for either in clinical practice. Moshina et al. suggest that pressure-standardised paddles may reduce image quality and increase AGD, although this requires additional research beyond the scope of this review to validate.

**Recommendations**

Current literature offers a pressure-standardised compression protocol of approximately 10kPa, with some variation. Although many studies recorded higher average values in the actual results, this can be attributed to subjective force standardisation as the foundation of data collection. One study indicated positive outcomes occur with a pressure reduction from 12–9kPa, another proposed an ideal force of at least 130N with associated pressure less than 9.8kPa, another recommends a standard pressure between 9.6 and 12kPa, and yet others suggest 10kPa results in an ideal tissue pressure between that of normal venous and arterial blood pressure. Research conducted by de Groot et al. similarly concluded that a compression of 10–12kPa corresponds with breast arterial pressure, although an increase in thickness of 9% for small and 2% for large breasts was also recorded. Whilst not ideal, this increase in thickness may be considered negligible due to the large dynamic range of digital mammography systems, as discussed previously. This notion is supported by Lau et al; a reduction in compression force of 32.5% has minimal effect on image quality and AGD. Low pressure and high breast volume are considered ideal for enhanced cancer detection, with pressures between 9.2 and 10.7kPa resulting in the highest detection rate. Further to this, high compression has been shown to reduce lesion visibility to a greater extent than insufficient compression. Mercer et al. also evaluate this from a mammographer’s perspective, suggesting that extending mammographer training may aid performance and technique consistency. Another recommendation includes the need for peer observation to allow for comparison of a mammographer’s own force compression practice to that of colleagues.

**Limitations**

Several drawbacks arose when analysing the literature, many of which are applicable across multiple studies. Firstly, generalising conclusions to an external population is unwise, as screening policies and populations differ across countries. This also raises the issue of extending results between screening and diagnostic mammography; as suggested by the literature, a diagnostic or post-intervention cohort may experience increased sensitivity to pain and hence are less tolerant to compression. A study conducted by de Groot et al. supports this concept; the odds for severe pain in
women post-breast-conserving therapy was 5.3 times higher than normal, largely due to changes in breast composition (elasticity).\textsuperscript{6,12} Accounting for differences between standard mammographic views (CC and MLO) is also important; the inherent discrepancies resulting from alternating the positioning technique limit the comparability of recorded values, both within and between studies.\textsuperscript{6,12} Additionally, the impact of pectoral muscle inclusion in the MLO contact area was reported; due to its presence, the computed pressure does not accurately reflect the pressure exerted on the breast tissue.\textsuperscript{6} Clinically, the appearance of specific lesion types experiencing over- or under-compression is inconclusive.\textsuperscript{6} Furthermore, variations across study types are useful to note; parameters available for acquisition and analysis differ depending on whether data collection was prospective or retrospective.

Limitations inherent in the review process itself must also be acknowledged. Timeframe limits were not applied, although dates were consciously noted throughout analysis; most studies were relatively recent regardless (i.e. all 2013 onwards, excluding two studies), given this field is currently evolving. Applying language-limiting parameters resulted in the omission of non-English sources; hence, scholarly evidence in foreign languages is not captured in this review. Upon analysis, most information originated from the Netherlands; this may demonstrate selection bias in that evidence for favoured outcomes is strengthened, yet similar conclusions were reached in the other literature. An extended search of more databases and ‘grey’ literature would confirm this. In fact, the distribution of resource origins serves to highlight the identified lack of Australian evidence, whilst reinforcing this review’s objective of pressure-standardisation protocol in a national context. Although potentially indicative of publication bias, this could not be assessed as no accepted method exists for its evaluation of diagnostic test accuracy studies.\textsuperscript{27} Furthermore, the selection of sources may display potential subjectivity as only two reviewers were involved in the process; reproducible criteria were adhered to in order to minimise this. Similarly, it is possible that data were missed due to keyword specifications and terminology issues.

**Conclusion**

The notion of mechanical standardisation techniques for breast compression is central to recent mammographic advances, with existing literature strongly supporting its implementation. Based on the current data available, several conclusions can be deduced. Key mammographic measures are obtained subjectively; the resultant disparity thus infers that optimal compression force values are not present in published research guidelines. Research supports an alternate standardised compression protocol founded on pressure; this approach accounts for individual breast characteristics in a ‘personalised’ manner.\textsuperscript{7,24} As well as an objective measure, a compression pressure of approximately 10kPa was found to decrease pain, with a negligible effect on breast thickness, AGD and resultant image quality.\textsuperscript{1,6–8,10,13,14,24} This also aids the reproducibility of image acquisition between and within women, whilst offering suggested guidelines for mammographers.\textsuperscript{7,10,13} It is therefore suggested that patient compliance would increase in accordance with the perceived benefits of a standardised technique, which would ultimately aid in the detection of early-stage breast cancer.\textsuperscript{9,16,17} However, recent literature also presents several limitations which may ideally be considered before widespread clinical application. This is particularly relevant as current Australian data are lacking; hence, the generalisation of evidence-based conclusions to a specific population requires additional research.

**Conflict of Interest**

The authors declare no conflict of interest.

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