Abstract

Purpose: To analyse whether the screening performance parameters of the Maltese National Breast Screening Programme first screening round met requirements set by European standards. The association between screening age and results of screening performance parameters was also investigated.

Method: Quantitative methodology was used to review examinations of women who were recalled for a technical recall or further assessment rates. All accessible members of the population recalled during the first round were retrospectively reviewed resulting in a sample of 2300 recalled examinations.

Results: Malta’s first screening round met the European Guidelines recommendations for technical repeat rate (0.26%), early recall rate (0.45%), breast cancer detection rate (13.77 per 1000 women) and Positive Predictive Value of screening test (7.58%). However, local recall rate (18.53%) and further assessment rate (18.27%) were higher than recommended.

The Chi square test showed a statistically significant difference ($p<0.05$) in recall rates between the compared age groups, as younger women (51-55 years) were more likely to have a negative diagnosis after the initial mammogram whereas older women (56-60 years) were more likely to be recalled. There was no age discrepancy ($p\geq0.05$) in local breast cancer detection rate and positive predictive value of screening test.

Conclusion: Although the Maltese first screening round performed well, this study found deficiencies in recall and further assessment rates, which according to literature may result in psychological morbidity and inefficient use of screening resources. This study also concluded that when a cohort is analysed, age is not as significant as the screening round itself (first/subsequent).

Keywords
Breast cancer, breast screening programme, screening performance parameter, false positive mammogram, age effect
Introduction

Breast cancer is the most common cancer among Maltese women and accounts for a third of all female cancer deaths. Every one in twelve Maltese women will get breast cancer in her lifetime; a significantly higher proportion than the European average. In October 2009, Malta’s Department for Health agreed to implement a National Breast Screening Programme (MNBSP). The aim of screening was to reduce morbidity and mortality from the disease without adversely affecting the health of participants. Key to achieving this aim were high levels of quality within the entire screening process. The MNBSP strives to protect the dignity and privacy of women, while offering an effective service at the highest levels of quality to diagnose and treat breast cancers at the earliest possible stage, and also meeting the European clinical standards.

The first MNBSP round commenced in October 2009 and ended in February 2013. The local screening programme provided free screening, every three years for all women aged 50 to 60, resident in the Maltese Islands. This age range was selected for the first screening round since this age group was deemed to be the ‘most at risk’ of developing breast cancer. Malta’s Department for Health additionally states that as soon as more human resources became available, the programme will be extended even to older women.

Double reading of mammograms was performed locally, as recommended for programmes in their first round of screening. When comparing dual to single reading, studies have shown that sensitivity increased by 5–10% without a significant effect on recall rates. Consensus arbitration of discordant double read cases also improved reading performance since it decreased recall rates and increased Positive Predictive Value (PPV). Since consensus arbitration was not practised locally, as per radiologists’ choice, women who were recalled by one radiologist experienced additional examinations during further assessment clinics. This could have resulted in higher recall rates and lower PPV.

A high quality screening service can be achieved through the use of targets, performance parameters and audits. Although audits of performance parameters were undertaken in European screening units, this research was the first of its kind in Malta. The findings offer valuable contribution for future advancement of the local screening unit as they helped to identify areas of strength, as well as areas needing improvement.
This study aimed to retrospectively audit results of the first screening round of the MNBSP and to assess whether the service fulfilled the European standard requirements. Thus the study objectives were to measure screening performance parameters (Recall Rate, Technical Recall (TR) Rate, Further Assessment Rate, Early Recall (ER) Rate, Breast Cancer Detection Rate, PPV of Screening Test) of the local first screening round; calculate the different types of clinical examinations performed during further assessment clinics; compare the rates of the screening performance parameters to the levels set by European and United Kingdom (UK) guidelines and investigate any association between screening age and results of screening performance parameters.

**Literature Review**

Several online databases such as Medline, Cumulative Index Nursing and Allied Health Literature, and ScienceDirect were utilised to access e-journals. Although most of the identified literature was undertaken after the inception of screening programmes, around 10–20 years ago, it was seminal to the study and was therefore included in this review. Only peer-reviewed, European studies written in the English language were included due to the similarity of breast screening programmes.

**Quality Assurance (QA)**

Ensuring the quality of a screening service is vital. This could be achieved by early monitoring of screening performance parameters of the unit, potentially optimising the use of resources and ultimately producing an observable reduction in mortality.\(^4\) Strict adherence to quality assurance and quality control guidelines must be practised in all mammography facilities to ensure accurate diagnosis, thus minimising false positive mammograms.\(^8,9\) False-positive rate refers to recalls for further assessment which turn out to be normal or benign.\(^10\) This is one of the reasons for the ongoing screening debate, since it gives rise to negative effects, namely: financial costs to the health service and psychological strain on the women.\(^10,11\) Since the majority of screening mammograms are normal, radiologists’ record of reporting should demonstrate high specificity avoiding false-positive mammograms.\(^12\)

In addition to false positive rates, several other performance parameters were identified by ‘The European Guidelines for QA in Breast Cancer Screening and Diagnosis’.
Recommended acceptable levels for each parameter were also set.\textsuperscript{4} Table 1 defines the screening performance parameters audited in this study and indicates the acceptable EU and UK levels for programmes in their first screening round.\textsuperscript{1,4,13,14,15}

Several research studies investigated these screening performance parameters.

**Recall Rate**

No increase in cancer detection rates and in screening sensitivity beyond a recall rate of 4.8% were recorded.\textsuperscript{16} High recall rates were found to signify that resources are used inefficiently in women undergoing unnecessary follow-up procedures.\textsuperscript{17,18} Conversely, rates lower than 1% were associated with reduced cancer detection and increased interval cancers.\textsuperscript{4} Previous research states that recall rates were influenced by several factors including training and experience of radiologists, image quality, the volume of mammograms interpreted and the age of screened women.\textsuperscript{19,20,21}

**Early Recall Rate**

ER was found to be associated with a low predictive value for malignancy and thus every effort should be made to obtain a definitive diagnosis at initial assessment.\textsuperscript{4,22,23,24} Additionally, a study analysing 110 women who were recalled early, revealed that 3.6% had invasive cancer, 0.9% had DCIS, while 84% had benign findings.\textsuperscript{25}

**PPV of Screening Tests**

The European guidelines did not define an acceptable PPV of Screening Test, whilst the UK National Health Service Breast Screening Programme (NHSBSP) defined the minimum standard PPV as 2.7% or more at first screenings.\textsuperscript{26} Due to the similarity between the UK NHSBSP and the MNBSP, this value was taken and accepted as the recommended level.\textsuperscript{27} Although recall rates of first screening rounds are generally high, the PPVs are usually low. In subsequent screenings the previous investigations could be referred to without the need for recall. Therefore only new findings would be fully investigated, thus lowering recall rates and further increasing PPV.\textsuperscript{28,20}

**Performance in European Breast Screening Programmes**

Performance indicators for mammography screening in 17 European countries showed some discrepancies, with recall rates ranging from 1.3% to 18.4%.\textsuperscript{29} First screening rounds resulted
in detection rates varying from 10.7 per 1000 women screened in Copenhagen to 3.6 per 1000 in Finland. This difference between countries should be interpreted with caution due to variations in screening and interpretation methods used within the various programmes. For instance the UK NHSBSP and the Netherlands followed very different recall policies; the latter deliberately aiming at a very low recall rate. Other differences included the use of one versus two-view mammography, screening interval, double-reading and methods for arriving at a resolution when double-reading led to different conclusions.

**Effect of Age**

Age is another independent factor predicting the accuracy of screening mammography. As screening age increased, recall rates decreased. Sensitivity, specificity and PPV increased with age thus increasing the accuracy of screening.

A study that analysed 215,665 mammograms revealed that the PPV was inversely related to the recall rates for age. Recall rates decreased from 7.3% for the youngest women to 4.9% for the oldest women, whilst the PPV rose from 1.9% to 12.7%. This outcome was supported by an extensive study of 1.5 million examinations reporting an increase in PPV by age. This was reasonable given the relatively higher breast density and lower incidence of disease in younger women.

Alternatively, studies analysing cohorts of women undergoing successive screening rounds indicated no clear association between age and risks of false-positive mammograms, concluding that such risks are much higher in first screening rounds. Nevertheless, there were no considerable reductions from second rounds onwards, regardless of women’s increasing age. Therefore, the association between age and false-positive recall rates seen in several cross-sectional studies could be overestimated because of the proportion of young women without a previous mammogram in first screening rounds. This implies that age is not as significant as the first screening round itself.

**Psychological Impact of Recall**

One of the reasons for maintaining screening performance parameters within the acceptable EU/UK levels is that women undergoing further assessment suffer significantly greater
adverse psychological consequences when compared to those with an immediate normal result. One study found that 30% of recalled women felt very anxious on receipt of their recall letter, whilst another study reported that 40% of women reported feeling very frightened when they received their recall letter. 16% of respondents felt certain that they had breast cancer. Such a psychological impact was reported by recalled women before, during, and immediately after recall. Women who had clear results after an invasive investigation and those who were placed on ER experienced the highest levels of anxiety.

The adverse psychological impacts of false-positive recalls could discourage women from re-attending for subsequent screening. Absenting from screening consequently negates any benefits of the service and increases the mortality rate. To ensure accurate and low false-positive recall rates it is important to regularly measure screening performance parameters and provide feedback to screening personnel.

Methods

Following ethical approval, a quantitative, non-experimental, descriptive, comparative, retrospective design was used to audit the examinations of women who were recalled for either a TR or further assessment. The study was conducted at the Maltese screening unit on the basis of data obtained from computerised databases. After every recall session, radiographers are responsible for recording all pertinent data on a database at the unit. This included recommended and performed TR and further assessments and outcomes of the recall clinics. All the women recalled during the first screening round (October 2009 to February 2013) were included in this study (n=2300). Retrospective data collection was performed between the 4th and 30th August 2014.

The information was audited by means of three data record sheets derived from the 4th edition of the European guidelines for QA in breast cancer screening and diagnosis. To provide a more sensitive measure of performance and to be able to investigate the effect of age on each parameter, data was further classified in age groups (51 to 55 and 56 to 60 years).

Data Record Sheet 1 (Table 2) was used to audit the possible outcomes of the initial screening mammogram, namely Negative Examinations, Radiologist Recommended TR,
Actually Performed TR, Radiologist Recommended Further Assessment and Actually Performed Further Assessment.

Data Record Sheet 2 (Table 3) was utilised to audit investigations performed during further assessment clinics, namely the Number of Clinical Breast Examinations, the Number of Additional Imaging, which was subdivided into Number of Further Views and Ultrasounds, and Number of Interventional Procedures, which was subdivided into Number of Radiologist Recommended Cytology and Actually Performed Cytology and Number of Radiologist Recommended WCNB and Actually Performed WCNB.

Data Record Sheet 3 (Table 4) was used to audit the outcome of the overall screening process, namely the Total Negative Examinations, ER following Further Assessment, Number of DCIS cases, Number of Invasive Cancers detected and Unknown Outcome. Data Record Sheet 3 was also used to audit the timing of breast cancer detection, namely At Initial Screen and During ER.

The researcher undertook the following process for data collection:

Recalled women were classified either as recommended TR or as recommended recall for further assessment. Such classification was opted for, for the sake of consistency with terminology used at the unit. The number of recommended and performed TRs and further assessments were audited. Data from each recalled woman was then reviewed. This served to audit: the number of women who had a clinical breast examination, the number of women who had additional imaging (further views or ultrasound); the number of women who underwent an interventional procedure (FNA or WCNB).

The number of women recalled for further assessment and classified as requiring an ER was also audited. Cases that had undergone interventional procedures were reviewed to determine whether any malignancy had been diagnosed. The type of cancer diagnosed was also documented and diagnosed cancers were audited according to the time of their detection (at the initial screen or during ER).

Descriptive statistics were used to present the audit results and to calculate screening performance parameters for the MNBSF. The chi-square test was used to establish whether
there was an association between categorical variables. One of the categorical variables was the age group (51-55; 56-60) whilst others described the screening performance parameters. The difference between two proportions test was used to determine whether local screening performance parameters that failed to meet acceptable levels set by European guidelines, differed significantly from the guidelines.

Results

The audit results and rates of screening performance parameter rates of the Maltese first screening round are summarised in Table 2-5. The MNBSP met the recommendations given by European Guidelines for TR, ER and breast cancer detection rates which were found to be 0.26%, 0.45%, 13.77 per 1000 women respectively. The PPV of the Screening Test for the Maltese first screening round was 7.58%, thus within the recommendations by the UK NHSBSP. However, local recall and further assessment rates (18.53% and 18.27% respectively) were higher than those recommended by European guidelines (Table 6).

The majority of recalled women (n=2251, 83.3%) underwent additional imaging (further views or ultrasound); whilst only 23 (0.9%) were examined clinically, 427 (15.8%) women underwent interventional procedures (FNA and WCNB). A total of 27 (0.2%) women were diagnosed with DCIS, whilst 144 (1.2%) women were diagnosed with invasive breast cancer. The majority of cancers (n=169, 98.8%) were detected at the initial screen whilst 2 (1.2%) cancers were detected in women for whom on ER assessment was recommended at further assessment.

The Chi square test showed a statistically significant difference (p<0.05) in recall rates between the compared age groups, as younger women (51 to 55 years) were more likely to have a negative diagnosis after the initial mammogram, whereas older women (56 to 60 years) were more likely to be recalled. There was no age discrepancy (p≥0.05) in the local breast cancer detection rate and PPV of the screening test.

Discussion

Higher recall rates were expected for first screening rounds when compared to subsequent ones. The local Recall Rate for the first screening round was higher than the acceptable rate for first screening rounds and also higher than European recall rates. The different
screening and/or interpretation methods utilized locally may account for the differences in recall rates leading to issues of comparability of findings to other studies and European recall rates.

Differences in data collection may have also affected comparability of recall rates. Although this research did not investigate the effect of breast density on mammography performance, studies have shown that this factor could affect the performance parameters. Screening programmes recording breast density demonstrated a great variation in breast density classification affecting comparisons of screening results.

The presence of symptoms at screening was another important modifier of performance estimates, as symptomatic women may have higher recall and PPV rates. A few programmes, including the local one, recorded symptoms present at the time of screening. This factor was not included in the pre-established data record sheets and was therefore not investigated in this research.

Whilst low recall rates may decrease detection rates and increase incidence of subsequent interval cancers, high recall rates alternatively cause psychological morbidity and indicate inefficient use of screening resources.

Additionally, differences in recall rates may have been affected by factors which were beyond the researcher’s control, such as the prevalence of cancer in the population, radiologists’ training, image quality, and in some instances, malpractice concerns. The training of radiologists is an important factor affecting the recall rate. The researcher had no evidence whether this factor affected the recall rate locally. However, the fact that this was the first experience for local radiologists to work within a screening program might have had an impact on the recall rate as acknowledged in literature. Local radiologists might have also recalled women with suspicious findings, even in the absence of a possibility of cancer, to avoid possible malpractice litigation. Additionally, reading an average of 3600 mammograms yearly, local radiologists are considered to be low volume readers compared with the stipulated 5000 yearly mammograms which European guidelines recommend. As this was the first unit on the island using direct digital mammography, radiologists were new to the equipment possibly leading to increased recall rates. Such high recall rates may also arise from the local absence of consensus or arbitration of double reading. Other studies have
suggested that double reading by consensus or arbitration, although being more labour intensive, decreases the recall rate.\textsuperscript{7}

The local Further Assessment Rate exceeded European guidelines for the same reasons since the recall rate is the summation of further assessment and TR rates.\textsuperscript{13} The small number of invasive investigations performed locally is considered as a strength since women with benign breast disease undergoing invasive investigations report higher levels of stress and anxiety lasting for several months after the additional investigations.\textsuperscript{43,50}

A TR rate of 0.26\% indicated high image quality locally, as the objective of a screening programme is to minimise the number of women undergoing technical recalls while achieving the optimum image quality.\textsuperscript{20} To obtain such an excellent TR Rate careful selection and training of staff is necessary.

Low predictive value for malignancy in ER has been expressed in various studies\textsuperscript{24,25} being consistent with this study’s findings as the majority of cancers were detected at initial screen, with only 2 cancers detected at ER stage. Nevertheless the latter could represent late diagnosis and failure of the screening and assessment process since women were falsely reassured, and ultimately diagnosed with cancer.\textsuperscript{4,51}

Such a promising Breast Cancer Detection Rate could be due to the benefits of direct digital mammography when compared to film screen mammography, namely improving detection of calcification related malignancies and non-symptomatic cancers. Although European guidelines do not specify an upper limit on the desirable levels, a high breast cancer detection rate may suggest a longer sojourn time (the period in which the tumour is asymptomatic but detectable by screening) than expected and diagnosis of cancers that would not have been detected without screening.\textsuperscript{52} However, over diagnosis has to be evaluated carefully with tumour characteristics in mind.

A PPV of the Screening Test of 7.58\% can be interpreted as 13 women recalled per cancer detected. Thus 13 women had a false positive mammogram for each cancer detected during the local first screening round. Studies have shown that false positive mammograms negatively affect women’s well-being; experiencing breast cancer related anxiety, worry and distress, when compared with women who receive normal mammogram results.\textsuperscript{10,11} Whilst
false positive mammograms are undesirable, they are necessary to maximise the number of small cancers detected.

Literature demonstrated that the age of the screened population has an impact on recall rates documenting a decline in rates across age and in subsequent screens.\textsuperscript{21,22} In contrast this research showed lower recall rates in the 51 to 55 years age group. Moreover, results revealed that younger women were more likely to have a negative diagnosis after the initial mammogram, and older women were more likely to be recalled. In this research the age of the screened population was 50 to 60 years, whilst in literature the age of the screened population was of a wider age span. However, comparing local results to other studies with wider age span was still appropriate.

This may be due to the fact that the programme commenced with the older client spectrum, and invited those aged 56 to 60 in the first years and then moved down the age group and invited women aged between 51 and 55 in the following years. Thus the reduction in recall rates over the years can be attributed to accumulated experience in screening, allowing individual radiologists to attain a greater level of experience and competence. However there was no age discrepancy in the local cancer detection rate and PPV of the screening test, signifying that mammograms had similar accuracy in both younger and older women.

**Limitations**

Limitations included the lacking or varying definitions of recall found in literature. Definition evidently affects the recall rate and thereby its relationship with PPV and breast cancer detection rate, making comparison of findings difficult. The performance parameters measured in this research were based on definitions stated by the European Guidelines\textsuperscript{4}, thus permitting comparison of findings to acceptable levels set by these guidelines.

Although European Guidelines were much more comprehensive, measuring additional screening performance parameters was impractical due to timing restrictions and available resources.
Participant related factors such as breast density and the use of HRT were a further limitation. These were not utilised for analysis and could have influenced screening performance parameters results. Further studies are required to elucidate this issue.

**Conclusions**

Through analysis of the first round of the newly implemented MNBS, it was concluded that the programme performed well since most of the performance parameters complied with European guidelines. This successful performance may be the result of joint efforts from all personnel, comprehensive QA measures and the use of digital techniques. Such positive performance can be used to build and maintain a high state of confidence among local radiologists and those responsible for the execution of the screening programme. A successful performance in the first screening round encourages new and repeat attendance to the Maltese screening programme.

This research has established that the local recall and further assessment rates did not comply with the acceptable levels set by the European guidelines, uncovering a problem in local clinical performance since according to literature high recall rates can cause psychological morbidity and indicating inefficient use of screening resources. In attempt to reduce recall and further assessment rates, the ongoing education of personnel particularly screening radiologists should be assured. Other recommendations include double reading by consensus or arbitration and interpretation of a high volume of mammograms.

This study also investigated any association between age and screening performance parameters and demonstrated that younger women were more likely to have a negative diagnosis after the initial mammogram whereas older women were more likely to be recalled. In addition there was no age discrepancy in the local cancer detection rate and PPV of the screening test, signifying that mammograms had similar accuracy in both younger and older women. Based on these outcomes, it was established that when a cohort is analysed, age is not as significant as the screening round itself.

Screening performance results highlight the importance of continuous surveillance of the local screening unit in order to sustain and improve performance and effectiveness. Thus it is suggested that regular QA is maintained to offer high quality mammographic screening.
Future research focusing on the effects of mammography equipment used, training of radiologists, volumes of mammograms read, and double-reading by arbitration is recommended.

**Conflict of Interest Statement**

Conflicts of Interest: None

**3803 Words**

**References**


Table 1: Screening Performance Parameters Definitions and Acceptable Levels for Programmes in their First Screening Round

<table>
<thead>
<tr>
<th>Screening Performance Parameters</th>
<th>Definition</th>
<th>Acceptable EU/UK Level</th>
</tr>
</thead>
<tbody>
<tr>
<td>Recall Rate</td>
<td>The number of women who have to physically return to the screening unit for a technical recall or for further assessment as a proportion of all women who had a screening examination.</td>
<td>&lt; 7</td>
</tr>
<tr>
<td>Technical Recall Rate</td>
<td>The number of women who have a repeat mammogram because of a technical inadequacy of the screening mammogram as a proportion of all women who had a screening examination.</td>
<td>&lt; 3</td>
</tr>
<tr>
<td>Further Assessment Rate</td>
<td>The number of women undergoing additional diagnostic techniques (clinical breast examination, additional imaging and invasive investigations) to clarify the nature of a perceived abnormality detected at the screening examination, as a proportion of all women who had a screening examination.</td>
<td>&lt; 7</td>
</tr>
<tr>
<td>Early Recall Rate</td>
<td>The number of women who are recalled for further assessment at an interval shorter than the normal interval (locally 3 years) as a proportion of all women who had a screening examination. Early recalls are performed when the initial further assessment examination leads to an equivocal diagnosis.</td>
<td>&lt; 1</td>
</tr>
<tr>
<td>Breast Cancer Detection Rate</td>
<td>The number of in situ and invasive malignant breast lesions detected in a screening round per 1000 women.</td>
<td>≥2.16</td>
</tr>
<tr>
<td>Positive Predictive Value of Screening Test</td>
<td>The number of cancers detected as a proportion of the women undergoing further assessment (excluding technical recalls).</td>
<td>&gt; 2.7</td>
</tr>
</tbody>
</table>
Table 2: Data Record Sheet 1 – Outcome of the Initial Screening Mammogram

<table>
<thead>
<tr>
<th>Age Group</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>51 to 55 years</td>
</tr>
<tr>
<td>Number of Women Screened</td>
<td>6781</td>
</tr>
<tr>
<td>Outcome of the Screening test</td>
<td></td>
</tr>
<tr>
<td>Negative</td>
<td>5623</td>
</tr>
<tr>
<td>Radiologist Recommended Technical Recalls</td>
<td>9</td>
</tr>
<tr>
<td>Actually Performed Technical Recalls</td>
<td>9</td>
</tr>
<tr>
<td>Radiologist Recommended Further Assessment</td>
<td>1149</td>
</tr>
<tr>
<td>Actually Performed Further Assessment</td>
<td>1143</td>
</tr>
</tbody>
</table>

Table 3: Data Record Sheet 2 – Investigations performed during Further Assessment Clinics

<table>
<thead>
<tr>
<th>Age Group</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>51 to 55 years</td>
</tr>
<tr>
<td>Number of additional imaging</td>
<td>1142</td>
</tr>
<tr>
<td>Types of additional imaging</td>
<td></td>
</tr>
<tr>
<td>Further views</td>
<td>907</td>
</tr>
<tr>
<td>Ultrasound</td>
<td>567</td>
</tr>
<tr>
<td>Clinical examination</td>
<td>8</td>
</tr>
<tr>
<td>Interventional procedures</td>
<td>213</td>
</tr>
<tr>
<td>Cytology</td>
<td></td>
</tr>
<tr>
<td>Radiologist Recommended Cytology</td>
<td>3</td>
</tr>
<tr>
<td>Actually Performed Cytology</td>
<td>3</td>
</tr>
<tr>
<td>Core Biopsy</td>
<td></td>
</tr>
<tr>
<td>Radiologist Recommended Core Biopsy</td>
<td>210</td>
</tr>
<tr>
<td>Actually Performed Core Biopsy</td>
<td>210</td>
</tr>
</tbody>
</table>
Table 4: Data Record Sheet 3 – Outcome of the Overall Screening Process and Timing of Breast Cancer Detection

<table>
<thead>
<tr>
<th>Age Group</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>51 to 55 years</td>
</tr>
<tr>
<td>Outcome of screening process</td>
<td></td>
</tr>
<tr>
<td>Negative Examinations</td>
<td>6678</td>
</tr>
<tr>
<td>Early Recalls following Further Assessment</td>
<td>28</td>
</tr>
<tr>
<td>Ductal Carcinoma in situ detected</td>
<td>10</td>
</tr>
<tr>
<td>Invasive Cancer Detected</td>
<td>87</td>
</tr>
<tr>
<td>Unknown Outcome</td>
<td>6</td>
</tr>
<tr>
<td>Timing of Breast Cancer Detection</td>
<td></td>
</tr>
<tr>
<td>At Initial Screen</td>
<td>95</td>
</tr>
<tr>
<td>At Early Recall</td>
<td>2</td>
</tr>
</tbody>
</table>

Table 5: Summary of screening performance parameter rates of the Maltese first screening round

<table>
<thead>
<tr>
<th>Screening Performance Parameters</th>
<th>51 to 55 years</th>
<th>56 to 60 years</th>
<th>Total Rates</th>
</tr>
</thead>
<tbody>
<tr>
<td>Recall Rate (%)</td>
<td>17.07</td>
<td>20.27</td>
<td>18.53</td>
</tr>
<tr>
<td>Technical Repeat Rate (%)</td>
<td>0.13</td>
<td>0.41</td>
<td>0.26</td>
</tr>
<tr>
<td>Further Assessment Rate (%)</td>
<td>16.94</td>
<td>19.86</td>
<td>18.27</td>
</tr>
<tr>
<td>Early Recall Rate (%)</td>
<td>0.41</td>
<td>0.50</td>
<td>0.45</td>
</tr>
<tr>
<td>Breast Cancer Detection Rate (per 1000 women)</td>
<td>14.30</td>
<td>13.13</td>
<td>13.77</td>
</tr>
<tr>
<td>Positive Predictive Value of Screening Test (%)</td>
<td>8.49</td>
<td>6.64</td>
<td>7.58</td>
</tr>
</tbody>
</table>
Table 6: MNBSP Screening Performance Parameters Results compared to Acceptable EU/UK Levels

<table>
<thead>
<tr>
<th>Screening Performance Parameters</th>
<th>MNBSP Results</th>
<th>Acceptable Level EU/UK Guidelines</th>
</tr>
</thead>
<tbody>
<tr>
<td>Recall Rate (%)</td>
<td>18.53</td>
<td>&lt; 7</td>
</tr>
<tr>
<td>Technical Repeat Rate (%)</td>
<td>0.26</td>
<td>&lt; 3</td>
</tr>
<tr>
<td>Further Assessment Rate (%)</td>
<td>18.27</td>
<td>&lt; 7</td>
</tr>
<tr>
<td>Early Recall Rate (%)</td>
<td>0.45</td>
<td>&lt; 1</td>
</tr>
<tr>
<td>Breast Cancer Detection Rate (per 1000 women)</td>
<td>13.77</td>
<td>≥2.16</td>
</tr>
<tr>
<td>Positive Predictive Value of Screening Test (%)</td>
<td>7.58</td>
<td>&gt; 2.7</td>
</tr>
</tbody>
</table>
Highlights

- The Maltese technical and early recall rates complied with European guidelines.
- Breast cancer detection rate and positive predictive value conformed to guidelines.
- The recall and further assessment rates exceeded recommended European guidelines.
- Effect of age on screening parameters is not as significant as the screening round.