Preventive Medicine and Medical Screening

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| One of flemain problems across the region is that we have scattered |
|---|
| campaigns instead of savening regularly |
| سجل موقوم من المحتوير بيجيني ١٠ - ٢٠ ألف أنتى مستان بعملوا وسنم |
| واحقال لما أعل السنة الجاي يرمعوا يجوا نخسهم (مسأهل هليون). |
| المقمم مم لدكتور إن هاي مم أكبر المستال في منطقة النزق المؤسط وهي إنه برل |
| ما يكوى دويدة مستى للغص المبر لعس طان النبى مثلًا ، بعل |
| "sattered campaigns". |
| ين إذي مؤتمرات دوعوين في مناطق مختلفة مرة ومدة أولفترة متسنة بالسنة و الي بحفرم |
| تَعَرَيْبًا ٢/ ٥٨٠كل ساء كمردن وبرج الشنى لسنة اللي تدها. |
| " هاي إنكرة (مؤحرات) ناجحة في البرابات ولكم للسَّى في وقدا الحالي " |

وأكّد على فكرت عثال إين احقالية معترر ناس كؤتر ستواعث أو شافوا عنه إىلان كن أقل مرعد المعنور لو اجتهم دعوة خاصَّة أو مكلة. يزمع لمثال سرطان التي , كرم كل امرأة في كماردن (٤٠ - ٢) منت في كل لمحد فظات تكون الها فراية متساوية في إعبول على فحص فيبحر. Okay, How can I salve the problem in Jordan, when I have million badies ?? (Make aplan 2) Make an objective of screening af least 700,000 females 3) Set a fime interval (in this example, we will set 2 years)

(M) Increase the working time of mammagram centers, why? (لم 6 زم ازبر عدد ساعان / أيام على لم اكن اللي يتفحف ؟ * استقال عدد أكبر * إتلحة غرصة لقحص المناء الحاملان في وقت الحطلة. (يعني إذا كان عل لمركز أحد الحني أخليه السبت - إحني) (5) leap in mind we have 20 manograms in Jordon (6) so year 1 - 230,000 year 2 - 230,000 المرحسان عدد أنام لحل وعدد إلمانكان بطلح تقريبًا فحص ١٠ ساء في اليوم dro minu le and إلى ", ... V leulo

and to make sure they'll come, I'll call each woman or send a tit of I'll do pilot study to see whethis the best way to organize this Man in Jordan Pilotsfuly negities (15 طبقًا هذا ليروتزكول معلق في السوير وبريطانيا وختَّال على عكس منعلقة النشق اكأوسط اللي تخصوا خلال قترة مست مع مدف مليون تشخص يخر إن 16 أقتل هذف أطول فيمن النجاح ، لفكرة أسبط ٧ و تفمن لكل إلساء vosi) & has

Preventive Medicine

 Prevention was defined by Last as: pincidence close to zero
 "Actions aimed at eradicating, eliminating, or minimizing the impact of disease or disability, or if none of these is feasible, retarding the progress of disease and disability".

it is difficult to test a million ladies for breast cancer, to simplify it, we make screening for 2 years duration, so that everyday about 10 ladies will be tested. they all must have the same chance to be tested... equal chance to be invited.



Spectrum of health and disease with the main strategies for prevention at each level



Stages. Health

Asymptomatic

Symptemedic

Primary prevention

Secondary Prevention

Tertiary

معخشاريتحا riskfactors oriclo

12,91

Prople withstage O discuse like: prediabetes, subclinical hypothyroidism, stage O concer management الاتي الشحص في هاي (treatment, what are the المرحلة قبل ما تطلح اعلى investigations)

Prtients with type I diabetes, terminally ill concer pationts

-reduce complications Preduce cance'r reoccurance

Outcome (Disability) Death Healthy) Quaternary prevention

using the best ways to breat

adisease

and getting theright treatment منال مرتفن سكري قبل ما أغير الدواء تبعم creative kukpase renal failure tiol citins

The Dr. said that he prefers tertiary prevention over Palliafive care (Queletil alert), melm?

1- It is wider covers 2- Palliative care (reduce pain, disability)

Medical Screening

1.you called a 45 women with no symptoms for breast cancer test... this is called systematic screening.

2. 45 women came for tonsillitis to hospital, and the medical provider advised her to have a breast cancer test, but she came for something else, this is called opportunistic screening.

What is screening the person has no symptoms, they are not seeking medical help. "The <u>systematic application of a test</u> or enquiry, to identify individuals at sufficient risk of specific disorder to **benefit** from further investigation or direct preventive action, among persons who have not sought medical attention on account of symptoms of that disorder." Wald,2004

3.a 54 years old man with a history of type 2 diabetes, he is coming for hypelipidemia , and was recommended to have a type 2 diabetes test ... opportunitve screening.

> appertunistic > systematic - no symptoms Screening

defection (irondeficiency Extremely Imp. (malignancy), constipation

Aims of screening

Better prognosis/outcomes for individuals

• Protection of public from communicable diseases

Example of successful medical screening

 Mortality from breast cancer by year of death for selected age groups, England and Wales, 1971-99



Year of death

Opportunistic screening (case finding):

- Do screening for someone when he/she comes into
- contact with the health system for another reason
 Solution of the system for another reason
 Solution of the system for your overweight or obese patients when they come to your clinic
- ---- Refer women within age criteria for cervical or breast cancer screening

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Screening versus diagnosis

• Early detection: symptoms and signs

المعانس بتخاف تنتشخف م

- It is essential to work in both directions in parallel way:
- Start your screening programs

&

Invest in early detection at GPs and selected specialties
 & general population levels awareness.

Delay in presentation, diagnosis and treatment for Breast cancer patients in Jordan



Figure 1: Proportion of participants by patient's delay, diagnosis delay, and treatment delay

Abu-Helalah, M., Alshraideh, A. H., Al-Hanaqtah, M. T., Da'na, M. D., Al-Omari, A., & Mubaidin, R. (2016). Delay in presentation, diagnosis, and treatment for breast cancer patients in Jordan. *The breast journal*, *22*(2), 213-217.

Delay in presentation, diagnosis and treatment for colorecrtal cancer patients in Jordan



Fig1. Proportion of participants by patient's delay, diagnosis delay and treatment delay

Abu-Helalah, M. A., Alshraideh, H. A., Da'na, M., Al-Hanaqtah, M. T., Abuseif, A., Arqoob, K., & Ajaj, A. (2016). Delay in presentation, diagnosis and treatment for colorectal cancer patients in Jordan. *Journal of gastrointestinal cancer*, *47*(1), 36-46.

Criteria for screening

there is a screening done even for rare diseases.

1. The disease/condition is an important health problem:



1. The disease/condition is an important health problem: Ryind under momen ales Well-defined disorder

• Known epidemiology (megnifude of complicated)

اقرأواعم كأعراف

• Well-<u>understood</u> natural history

بوجوا مستون بالمعالي الطبيب معنان بروم بر بروجوا مربور المحتور بالمعنور بالممنور بالمامار بالمالمامار بالمالممنور بال Prevalence of undiagnosed cases

لوکان کیرهون بحتاج جمام

Shall we screen only for common illnesses?

- For serious diseases, even if it is not highly prevalent.
 e.g. <u>Neonatal</u> screening for inborn errors of metabolism.
- Phenylketonuria screened for in the UK. Incidence 1:12000 live births.
- If undetected, it would lead to severe mental retardation and growth retardation. While detected cases could be treated simply by dietary restriction of phenlylalanine.
- If undetected leads to severe mental and growth retardation. Early Detected cases easily treated by dietary restriction of PKU.

Congenital hypothyroidism screening in Jordan

* Thalasenia before marriage

2. Presence of presymptomatic or early

stage

- Is there an evidence from a randomised controlled trial that an earlier intervention would work?
- Detecting the disorder at this stage should help in getting better outcomes when compared with the situation without screening.
- Randomised controlled clinical trials could be needed to evaluate the impact of treatment on those detected from screening programmes as they could be different from those seeking medical attention for their conditions.
- Screening for a disease or a risk factor
 It is recommended to screen for diseases, while risk factors are bad screening tools

| Diabetes test | Normal | Prediabetes | Diabetes |
|--------------------------------|--------|-------------|----------|
| Hemoglobin A _{1C} , % | < 5.7 | 5.7-6.4 | ≥ 6.5 |
| Fasting blood glucose, mg/dL | < 100 | 100-125 | > 125 |
| Oral glucose tolerance, mg/dL | < 140 | 140-199 | > 199 |

| Trial | Design | Subjects | N; duration (years) | Control group | Active treatments | % change in diabetes risk |
|--|-------------|-------------------------------------|---------------------------|---|---|-----------------------------------|
| Principal diabetes preven | tion trials | s that evaluated metfor | rmin | | | |
| DPP (US) [19] | RCT | IGT and high- normal glucose | 3234; 3 | Placebo plus standard lifestyle advice | Metformin plus standard lifestyle advice | -31 -58 |
| DPP Outcome Study (US) [69] | 0 | Epidemiological follow-up to DPP | 2766; 5.7 | Placebo plus intensive lifestyle advice | Metformin 1700 mg/day + intensive lifestyle advice | -13 +5 |
| IDPP (India) [20, 65] | RCT | IGT | 531; 2.5 | Standard lifestyle advice | Intensive lifestyle advice Metformin plus standard lifestyle advice | -26 -28 |
| | | | | | lifestyle intervention Intensive lifestyle intervention | -29 |
| Wenying et al. (China) [68] | NR | IGT | 321; 3 | Standard lifestyle advice | Metformin Acarbose Intensive lifestyle intervention | -88 -87 -43 |
| Li et al. (China) [66] Iqbal Hydrie et al. (Pakistan) [67] | RCT RCT | IGT IGT | 70; 1 317; 1.5 | Placebo Standard lifestyle advice | Metformin Metformin Intensive lifestyle intervention | -66^{a} -76.5 |
| CANOE (Canada) [64] | RCT | IGT | 207; 3.9 | Placebo | Metformin 500 mg plus rosiglitazone 2 mg twice daily | -66 |
| Principal diabetes preven | tion trials | s that did not evaluate | metformin | | | |
| Diabetes Prevention Study (Finland) [70] | RCT | IGT | 522; 3.2 | Standard lifestyle advice | Intensive, multifactorial lifestyle intervention | -58 |
| Da Qing study (China) [71] | RBS | IGT | 577; 6 | Standard lifestyle advice | Diet, exercise, or both together | -31 to -46 |
| STOP-NIDDM (International ^b) [72, 73] | RCT | IGT | 1429; 3.3 | Placebo | Acarbose | -25 |
| XENDOS (Sween) [74] | RCT | IGT and obesity | 694; 4 ^c | Placebo | Orlistat | -45 |
| DREAM (21 countries ^d) [75, 76] | RCT | $IGT \pm IFG$ | 5269; 3 | Placebo Placebo | Rosiglitazone Ramipril | -62^{e} -9 ^f (NS) |
| IDPP-2 (India) [77] | NR^{f} | IGT | 407; 3 | Placebo + lifestyle intervention | Pioglitazone + lifestyle intervention | +8 (NS) |
| SOS study (Sweden) [78] | RCT | Obese, non- diabetic | 3429; 10 | No surgery ^g | Bariatric surgery | -83 |

ORIGINAL ARTICLE

A randomized double-blind crossover trial to investigate the efficacy of screening for adult hypothyroidism

M Abu-Helalah, M R Law, J P Bestwick, J P Monson and N J Wald

J Med Screen 2010;17:164–169 DOI: 10.1258/jms.2010.010057

Objective To assess the value of population screening for adult hypothyroidism. **Setting** Healthy people attending for a general health assessment.

Methods A thyroid-stimulating hormone (TSH) measurement was performed on people attending for a general health assessment (women aged 50–79 [35–49 with a family history of thyroid disease] and men aged 65–79). Those with TSH levels above 4.0 mU/L were invited to join a randomized double-blind crossover trial of thyroxine and placebo, each given in random order for four months. On entry a second blood sample was collected for a TSH measurement after the end of the trial to determine whether this would help select individuals for thyroxine treatment. The daily thyroxine dose started at 50 µg and if necessary was increased to achieve a TSH level of 0.6–2.0 mU/L.

Results There were 341 (8%) people with a TSH level above 4.0 mU/L, 110 met eligibility criteria (64 agreed to participate), and 56 (49 women, 7 men) completed the trial. Among the 15 individuals with a repeat TSH measurement above 4.5 mU/L, 11 reported feeling better on thyroxine than placebo and none reported feeling better on placebo (P = 0.001; four felt no different), indicating that in this group 73% benefitted (i.e. 11/15; 95% CI 45–92%). The main symptoms relieved were tiredness and loss of memory. There was no indication of harm. In the 41 individuals with a repeat serum TSH of 4.5 mU/L or less: 10 reported feeling better on thyroxine than placebo and 16 better on placebo (P = 0.42, 15 felt no different). Thus about 8% of men and women in the specified age groups had a TSH above 4.0 mU/L, and of these about a quarter had a repeat TSH above 4.5 mU/L, of whom about half would benefit from thyroxine treatment.

Conclusion The results indicate that screening for hypothyroidism would be worthwhile. Approximately 1% of people screened would have a better quality of life. Pilot screening programmes for adult hypothyroidism are justified.

See end of article for authors' affiliations

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Accepted for publication 25 August 2010

What do you aim to achieve from your screening programme?

Mortality



Quality of life and psychological wellbeing

Screening test:

 Safe Inexpensive -> (colpteral concerscreening agreed test but too expensive, 500\$9 x Acceptable (عبت اختبار ورحبت عدة مرة ثانية = نفس النتيجة) « Reliable • (قد ب يقدر بحدد لمريف بشكل لمعيم) - Valid • No or minimal adverse effects: pain or any possible adverse effects should be considered in addition to convenience and duration of the test.

مرم جبرًا. Screening test validity

• The validity of a screening test can be evaluated through its detection rate (sensitivity) and specificity.

ability of your tool to find the disease.

A. Detection rate (sensitivity) evaluates the ability of a screening tool to detect the disorder or problem. It represents the proportion of <u>diseased</u> individuals who have a positive screening test.

B. Specificity is the ability of a screening tool to label people without the targeted condition as "unaffected" (for diseases, healthy people as non-diseased).

T+ label diabetic = diabetic sensitivity F+ label diabetic = healthy 1-size ceficity T-s label healthy = healthy speceficity Fs label healthy = diabetic 1-sensitivity

1 diabetric patients 7 non-diabetric

م موجدًا



An ideal laboratory test would detect all people who have a disease and at the same time identify as normal all those who do not have the disease



Test score

False positive rate (1-specificity)

- More meaningful and practical than specificity because it shows the expected rate of those who would be falsely labelled as diseased or screen positive and might offered further investigations.
- It helps in estimation the magnitude of the ecohomic (further investigations) and other harmful effect such as psychological distress associated such outcomes.

الأثار النفسة السينة بعداجية للتشخيص بخطئ ولما يتعل اختبار ناف لتأكير رج يكور مكلف Validity of a test

How well a test performs can be assessed based on the values in the following 2x2 table

| | Disease | Disease |
|----------------------------------|----------------------|--------------------|
| | present | absent |
| Test positive or | True Positives | False positives |
| Surveillance | TP | FP |
| Detection positive | a | b |
| Test negative or Surveillance | C False negatives | d True negative |
| Detection negative | FN | TN |

| | Disease | Disease | |
|----------------------------------|-----------------------------|--------------------|-----------|
| | present | absent | |
| Test positive or | True Positives | False positives | |
| Surveillance | ТР | FP | 2 |
| Detection positive | a | b | 1-Specif. |
| Test negative or Surveillance | C False negatives | d True negative | + TN |
| Detection negative | FN | TN | TN+FP |

| Sensitivity $=$ - | Diseased people with a positive test _ | TP |
|-------------------|--|---------|
| | Alldiseasedpeople | TP + FN |

| Specificity = | Well people with a negaitive test | |
|---------------|-----------------------------------|--------------------|
| | All well people | $\overline{TN+FP}$ |

False positive rate= FP/FP+TN




False positive rate

• The proportion of unaffected individuals with positive test results.

False positive rate= b =1-specificty b+d

Predictive values

onto evaluate

Not

- Positive predictive value= all true positives/all positives(all true and all false) ×100
- How likely it is that a positive test result indicates
 the presence of the disease.
- It is the percentage of all people who test positive and who really have the disease
- Negative predictive value= True negatives/all negatives ×100
- It is the percentage of all people who test negative who really do not have the disease

'AN EXAM QUESTION: *What is the vight best to assest performance by a screening test? I've predictive values. X How boassist the screen fragram? sensitivity + FP rate Gyou can also add specificity but Tare more imp.

| | Disease | Disease | | |
|--|-----------------------|-----------------------|--|--|
| | present | absent | | |
| Test positive or Surveillance | True Positives TP | False positives FP | | |
| Detection positive | a | b | | |
| Test negative or Surveillance Detection negative | c | d | | |
| | False negatives FN | True negative TN | | |

$$prevalence = \frac{Diseased \ people}{All \ people} = \frac{TP + FN}{TP + FN + FP + TN}$$

predictive value positive = $\frac{Diseased people with a positive test}{All people with a positive test} = \frac{TP}{TP + FP}$

predictive value negative = $\frac{Well \text{ people with a negative test}}{All \text{ people with a negative test}} = \frac{TN}{TN + FN}$

Screening test validity:

Outcomes of screening tests

| | Disease present | Disease absent | | All |
|---------------------------|--|------------------------------|-------------------------|-------------|
| Positive screening test | <i>a</i> (true positive) | <i>b</i> (false positive) | | a + b |
| Negative screening test | <i>c</i> (false negative) | <i>d</i> (true negative) | | c + d |
| All | a + c | | b+d | a+b+c+d |
| Detection rate | proportion of af individuals with po test results | fected ositive | <u>_a_</u> a+c | |
| Specificity | Proportion of unaf individuals with ne test result | fected gative | $\frac{d}{b+d}$ | |
| False positive rate | proportion of unaf individuals with po- test results | fected ositive | $\frac{b}{b+d} = (1-s)$ | pecificity) |
| Positive predictive value | Probability of the d being present giv positive test | lisease en a | $\frac{a}{a+b}$ | |
| Negative predictive value | probability of no d being present giv negative test result | lisease en a | $\frac{d}{c+d}$ | |

| | | Patients with bowel cancer (as confirmed on colonoscopy) | | |
|---|----------|--|---|---|
| | | Positive | Negative | |
| Fecal Occult Occult blood screen test outcome | Positive | True Positive (TP) = 20 | False Positive (FP) = 180 | $\rightarrow \frac{\text{Positive predictive value}}{= TP / (TP + FP)}$ $= 20 / (20 + 180)$ $= 20 / 200$ $= 10\%$ |
| | Negative | False Negative (FN) = 10 | True Negative (TN) = 1820 | → Negative predictive value = TN / (FN + TN) = 1820 / (10 + 1820) = 1820 / 1830 ≈ 99.5% |
| | | $\downarrow Sensitivity = TP / (TP + FN) = 20 / (20 + 10) = 20 / 30 \approx 66.67%$ | ↓ Specificity = TN / (FP + TN) = 1820 / (180 + 1820) = 1820 / 2000 = 91% | |

Example of validity assessment

| | G-FOBT | FIT | | |
|---------------------------|----------------------|----------------------|--|--|
| Sensitivity | 50.00% (6.76-93.24) | 75.00% (19.41–99.37) | | |
| Specificity | 77.87% (72.24-82.83) | 90.12% (85.76–93.50) | | |
| Positive likelihood ratio | 2.26 (0.83-6.18) | 7.59 (3.86–14.94) | | |
| Negative likelihood ratio | 0.64 (0.24-1.71) | 0.28 (0.05-1.52) | | |
| Positive predictive value | 3.45% (0.42-11.91) | 10.71% (2.27-28.23) | | |
| Negative predictive value | 98.99% (96.42-99.88) | 99.56% (97.59–99.99) | | |

False positive rates: 1-Specificity More un-necessary colonoscopes and more cost for the program

Reliability of screening test

- Reliability means that the same results should be obtained by different observer or the same observer at different occasions.
- In practice, it is hard to achieve 100% reliability
- Guidelines should be in place on decisions when
- two observers have different opinions.

Agreed plan on further investigation, diagnosis and treatment:

It's unethical to have people with a trefest (19891 in gliger 109

- Where to refer your positive subjects
- What is the diagnostic tests
- Who will pay for the investigations and treatments
- Diagnostic tools, screening intervals and treatment
- Facilities required for such steps should also be available or easily installed and equally accessed by the screened population

Systematic application

 This means that the test is offered routinely to the target group <u>based on agreed criteria</u>.

Do it in a systematic way!

- Regular systematic national screening programs for breast and colorectal cancers should replace the current scattered campaigns and activities in many countries in the region. (مافيداى المناء الجار لأنه كارم ا السرطان المكي المناء الجار لأنه كارم ا
- Work should start with pilot systematic screening projects in representative area in the country of interest.



Obtain data from Ministry of Interior on residents in Areas 1,2,3 who fulfills screening criteria

Send letters through Health Centers C1,C2,C3

Send reminders through Health Centers C1,C2,C3 for non-respondents

Ask practice manager or health counselor to call non-respondents from the two calls and arrange for GP visit if needed.

Obtain data from the screening centers for respondents to screening calls.

Simplify your program

- Is it too difficult to have a national systematic regular screening program for breast cancer in country "x" where the number of women aged 40-70 is 1,000,000?
- In this country: it is recommended to screen women aged 40-69 once every two years
- Notice: Screening interval depends on mean sojourn time and should not be fixed to be on annual basis unless there is clinical evidence for that

Cut it down so it will be simple

Practical example: In country X, there are 1000000 women aged 40-70 who are eligible for screening

| 100000 | Women aged 40-7 | 0 | | | | | |
|--|------------------------------------|---|-------------------|------------|--------|---------|----|
| To be screened ann | ually | 50000 |) | ``` | | | |
| | | | | \sum | r. Rea | ad It, | |
| 75% response rate: | | 375000 |) | C | | | ٨ |
| 300 working days/ | 6 days work | | 1250 | | o ne | ef it a | f. |
| if there are 12 main districts in your country | | | | | | | |
| 25 centers in the whole country | 2 mammograms per center | 50 mammogra | ns | | | | |
| 1250/50 | 25 subjects Per machine per day | 7 working hours, means 4 subjects per hour | In the UK, 6-8 pa | tients per | | | |
| If we have only 5 centers in Amman, 3 centers in Irbid, 2 centers in Zarqa, 2 centers in Karak and one center in the remaining governorates | | | | | | | |
| we need 50 machines in 25 centers for 1 million women across Jordan | | | | | | | |
| This number is already available and can be provided at the public sector | | | | | | | |
| | | | | | | | |

Breast self-examination and death from breast cancer: analysis

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Breast self-examination (BSE) is widely recommended for breast cancer prevention. Following recent controversy mammography, it may be seen as an alternative. We present a meta-analysis of the effect of regular BSE on breast From a search of the medical literature, 20 observational studies and three clinical trials were identified that reported death rates or rates of advanced breast cancer (a marker of death) according to BSE practice. A lower risk of mobreast cancer was only found in studies of women with breast cancer who reported practising BSE before d pooled relative risk 0.64, 95% CI 0.56–0.73; advanced cancer, pooled relative risk 0.60, 95% CI 0.46–0.80). The n due to bias and confounding. There was no difference in death rate in studies on women who detected their examination (pooled relative risk 0.90, 95% CI 0.72–1.12). None of the trials of BSE training (in which most practising it regularly) showed lower mortality in the BSE group (pooled relative risk 1.01, 95% CI 0.92–1.12). T BSE is associated with considerably more women seeking medical advice and having biopsies. Regular BSE is not a of reducing breast cancer mortality. *British Journal of Cancer* (2003) **88**, 1047–1053. doi:10.1038/sj.bjc.6600847 www.bjcancer.com

Keywords: breast self-examination; breast cancer; mortality; meta-analysis



Figure 1 Observational studies of women with breast cancer, comparing the breast cancer death rates between the BSE and non-BSE groups. A test for heterogeneity between the studies yielded a *P*-value of 0.41 for those studies based on women who practise BSE and a *P*-value of 0.26 for those based on finding cancer by BSE.











Test it before you generalize it

حكى جها وانادى فها الما-

- Start with pilot program : (response) 2/1/100)
- Assess response rate
- Is my program cost-effective
- What is my cost-effective screening criteria
- Quality of all involved steps (single versus double reader mammography screening, FIT versus Haemoccult test)
- Compare respondents with non-respondents
- Assess success rates
- Look for determinants of success and failure
- Is there a specific group who needs different intervention?

Importance of Pilot Projects

ilot studyis done offer assernant on the Population

- 1. Health economics evaluation
- 2. Setting age cut-off based on local data
- 3. Improve performance at national level by learning from experience at pilot phase
- 4. Comprehensive assessment of the screening program helpline, waiting time, film quality, guidelines such as double readers, false positive rate, false negative rate, diagnosis process, psychological counseling, treatment, prognosis, economic evaluation, how can we make it better at the national level.
- 5. Assessment of barriers to screening
- 6. Quality assessment of staff

Acceptability of programme to the public and health care staff.

 Screening test, diagnostic test and therapeutic options should be ethically and socially accepted by the general public and the health care professionals.

Economic evaluation:

- Implementing screening programmes should be more economically effective than the existing system.
- Cost of all steps related to the screening programme should be assessed and compared with outcomes of the screening and with other services.
- Each country should has its own studies and data
- What is cost effective in the UK might not be cost effective in Jordan or India
- In breast cancer screening: age range for screening plays a key role in the cost-effectiveness of the program
- UK (Screening aged 50-70 Every three years, then in few years ago aged 40-49 at high risk)
- Sweden (age 40-70) annually

FURCIPIAN FOURIAR OF PURE CHIALTH 1997. 1-68 76

MAMMOGRAPHIC SCREENING

Economic evaluation of a mammography-based breast cancer screening programme in Spain

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The aim of the study was to perform a cost-effectiveness analysis of a breast cancer (BC) mammography screening programme, compared to a do-nothing alternative, in Spain. Screening consisted of a biennial mammography performed on all women 50–65 years old. A marginal analysis including women 45–49 years old was also performed. With the aid of a decision tree model, the numbers of BC cases diagnosed through screening, BC cases missed by screening and false-positive BC cases were calculated. Costs were calculated by feeding local data into Markovian models and the cost-effectiveness ratio calculation was performed in a computer spread sheet. A sensitivity analysis was also conducted. Results were presented in ECUs of 1993. The cost-effectiveness ratio per avoided death is 115,500 ECUs and per saved life year 7,300 ECUs. Including women 45–49 years old in the programme raises this ratio to 229,000 and 9,400 ECUs respectively. The sensitivity analysis showed the efficacy of mammography, compliance of the programme and screening costs to be the more sensitive variables.

Key words: breast cancer, screening, economic analysis, cost-effectiveness analysis

Bias related to medical screening

• Lead time bias: screened cases are detected at an earlier stage than that in which treatment would be worthwhile.

Does treatment work better at this stage?

 Length time bias: cases detected through screening are slowly progressive and may not harm the patient in lifetime

Selection bias: respondents are different from decliners

Volunteer bias:

• They tend to be of higher socioeconomic class

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- More health-conscious
- Comply better with prescribed advice
- Therefore, better results for a screening programme of volunteers compared with disease outcomes for non-voluntees may be relate to factors associated with the "volunteerism" rather than benefits of treatment following diagnosis.
- Therefore it is essential to analyse data on participants and ensure that all target group have the same access and received the same message

early diagnosis of a disease falsely makes it look like people are surviving longer.

Lead time bias

- Lead time: period between when the disease is detected by screening and when it would have become symptomatic and been diagnosed in the usual way.
- Prolongation between diagnosis and death

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- There is no difference in outcomes between patients detected through screening and patients who is treated when the condition <u>manifest</u> clinically
- Screening simply makes the condition evident at an earlier stage without actually affecting its course. (appears to lead to longer survival because of earlier detection)
- If left with no screening the disease will be diagnosed at age of 50 and die at age of 54
- If screened disease will be diagnosed at age of 47 and die at the age of 54



Lead time bias in Prostate cancer

- Lead Times and Over detection Due to Prostate-Specific Antigen Screening: Estimates From the European Randomized Study of Screening for Prostate Cancer
- Gerrit Draisma Rob Boer Suzie J. Otto Ingrid W. van der CruijsenRonald A. M. Damhuis Fritz H. Schröder Harry J. de Koning
- JNCI: Journal of the National Cancer Institute, Volume 95, Issue 12, 18 June 2003, Pages 868– 878, <u>https://doi.org/10.1093/jnci/95.12.868</u>

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Length time bias

• It is a form of selection bias.

 When we screen for disease were more likely to detect cases where the disease is progressing slowly

(we don't do for levkemin for eq)

overestimation of survival duration

• Over-presentation of slowly progressing disease among cases detected by screening.

 Screening will detect more slowly growing tumours, while rapidly growing tumours are more likely to develop and to proceed to clinical presentation within the interval between two consecutive screening examinations.

lead time bias; lead to early detection... so there is a long duration and we make screening for them. length time bias; slow cases being detected , slowly progression, we don't make screening for them.

Length time bias

 Faster-growing <u>tumors</u> generally have a shorter asymptomatic phase than slower-growing tumours, and so are less likely to be detected. However, faster-growing tumors are also often associated with a poorer prognosis. Slowergrowing tumors are hence likely to be overrepresented in screening tests. This can mean screening tests are erroneously associated with improved survival, even if they have no actual effect on prognosis.



Prostate Cancers With Varying DPCPs



DPCPs: detectable preclinical phase

Challenges

- Validity of the screening test
- Healthy people need further tests
- Anxiety caused
- Health care resources

Pilot basis

• What is my next step?

Quality Assurance

- Quality assurance means that the assessment of the service provided and applying modifications when necessary.
- This includes various steps such as recruitment, registration, waiting time, test procedures, results handling and follow up or referral for treatment procedures.

Clinical audit

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My programme is already in place Continuous monitoring and regular evaluation

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