Chapter 12:

Sylvia K. Plevritis, Bronislava M. Sigal, Peter Salzman, Jarrett Rosenberg, Peter Glynn

Background: We present a simulation model that predicts U.S. breast cancer mortality trends from 1975 to 2000 and quantifies the impact of screening mammography and adjuvant therapy on these trends. This model was developed within the Cancer Intervention and Surveillance Network (CISNET) consortium. Method: A Monte Carlo simulation is developed to generate the life history of individual breast cancer patients by using CISNET base case inputs that describe the secular trend in breast cancer risk, dissemination patterns for screening mammography and adjuvant treatment, and death from causes other than breast cancer. The model generates the patient’s age, tumor size and stage at detection, mode of detection, age at death, and cause of death (breast cancer versus other) based in part on assumptions on the natural history of breast cancer. Outcomes from multiple birth cohorts are summarized in terms of breast cancer mortality rates by calendar year. Result: Predicted breast cancer mortality rates follow the general shape of U.S. breast cancer mortality rates from 1975 to 1995 but level off after 1995 as opposed to following an observed decline. Sensitivity analysis revealed that the impact adjuvant treatment may be underestimated given the lack of data on temporal variation in treatment efficacy. Conclusion: We developed a simulation model that uses CISNET base case inputs and closely, but not exactly, reproduces U.S. breast cancer mortality rates. Screening mammography and adjuvant therapy are shown to have both contributed to a decline in U.S. breast cancer mortality. [J Natl Cancer Inst Monogr 2006;36:86–95]
Model Overview

Figure 1 presents a flowchart of our simulation model. The model is a continuous-time Monte Carlo simulation that can generate the life histories of individual patients under various dissemination profiles of screening mammography and adjuvant treatment. Characteristics generated for a simulated individual breast cancer patient include the following: 1) the date of her birth; 2) the age of her death from causes other than breast cancer; 3) her primary tumor volume doubling time, tumor volume at clinical detection, and tumor volume at the transitions to regional and distant stages; 4) the ages that she undergoes mammographic screening examinations; 5) the ages that she is detected with invasive breast cancer in the absence and presence of screening; 6) her primary tumor volume, extent of nodal and distant involvement ER status at detection, in the presence and absence of screening; 7) the adjuvant treatment that she received, dependent on her tumor characteristics and disease stage at detection; 8) her breast cancer survival time given her disease stage, size, and age at detection; and 9) her cause of death (i.e., breast cancer versus other causes).

In brief, to predict outcomes of breast cancer patients in the presence and the absence of screening, we start by reconstructing the natural history of the patient’s disease; next, we superimpose a screening schedule onto the natural history model; and finally, we determine if and when the screening test interrupts the natural history. Our natural history model describes the tumor size and stage of invasive cancer during its preclinical course. The screening schedule specifies the patient’s age at the time of screening mammography. A patient is screen detected only if the size of her tumor is at or above the tumor size detection threshold of mammography at the time of screening. Fig. 2, A and 2, B illustrate the interaction between the natural history model and the screening test for a patient whose tumor is detected by mammography and whose tumor is detected because of symptoms, respectively. Once the patient is detected, she is assigned a breast cancer–specific survival time dependent on her age, tumor size, stage, mode of detection, and her use of adjuvant treatment. Her age of death is the minimum age of breast cancer death and the age of other-cause death. Individual-level outcomes are summarized as population level outcomes in terms of age-adjusted breast cancer incidence and mortality rates by calendar year.

Table 1 provides the pseudocode for our simulation algorithm. The code was developed in C/C++. To generate individual-level outcomes for approximately 100 birth cohorts with 2 million individuals per birth cohort, the algorithm runs for approximately 2.5 hours to run on a Sun Solaris machine.
We built our model by first identifying clinically and biologically reasonable components that would influence breast cancer mortality. Two examples of such components include the natural history of the disease and the detection characteristics of the screening test; all model components are described in “Model Components.” Many of the components rely on the use of CISNET base case inputs (3). To enable a direct comparison between our model and other models that were built in the CISNET Breast Cancer Consortium, we did not modify the CISNET base case inputs. Inputs unique to our model describe the natural history of breast cancer and the detection characteristics of screening. We formulated these model-specific inputs with a few parameters to ensure identifiability. We merged all components into a Monte Carlo simulation algorithm that generates the breast cancer outcomes for individuals across multiple birth cohorts. We evaluated our model by demonstrating how well it reproduces breast cancer mortality trends. Because all factors influencing breast cancer mortality are not known, we did not calibrate to breast cancer mortality in the model building process. We regard a reasonable difference between the predicted and observed mortality rates as a measure of the effect of other factors that we did not incorporate in our model because of the lack of informative data.

**Model Components**

Figure 1 identifies the underlying components of our simulation model and the input(s) to each component.

**Population Component**

Our population component specifies the birth cohorts underlying the U.S. breast cancer trends from 1975 to 2000. To reproduce the outcomes of women aged 30–84 years in each calendar year from 1975 to 2000, we generated a sample of U.S. women born between 1887 and 1970. Each birth cohort consists of 2 million women, which we found to be a large enough number to reduce the sampling variability of the Monte Carlo method. Even though factors such as population immigration and emigration are likely to vary the relative sizes of the birth cohorts in the real world, the size of each birth cohort is kept constant in our simulation model because the incidence and mortality trends are reported as age-adjusted rates. Each simulated woman is assigned a birth date and an age at death from other causes. Given the year of birth, the date is uniformly distributed within the year. Similarly, given integer age at death of other causes, actual age is simulated uniformly within the year. Death from breast cancer and other causes are assumed to be independent. This component relies on a CISNET base case input, which specifies the other-cause death rate, by birth cohort, based on the Berkeley Mortality Database (4).

**Breast Cancer Incidence Component**

Our breast cancer incidence component determines whether or not an individual from a particular birth cohort would become

<table>
<thead>
<tr>
<th>Table 1. Pseudocode of Monte Carlo simulation algorithm developed to predict U.S. breast cancer incidence and mortality rates from 1975 to 2000</th>
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| For birth cohorts 1887 to 1970  
For individuals 1 to 2000000  
Step 1: Generate date of birth (see Population Component)  
Step 2: Generate age at other-cause death given birth cohort (see Population Component)  
Step 3: Generate age at clinical detection (see Breast Cancer Incidence Component)  
Step 4: Generate tumor volume doubling time (see Natural History Component)  
Step 5: Generate tumor size at clinical detection given the tumor volume doubling time (see Natural History Component)  
Step 6: Generate tumor size at the onset of regional and distant stage (see Natural History Component)  
Step 7: Compute stage of the tumor at clinical detection  
Step 8: Generate age at breast cancer death following clinical detection given age and stage at clinical detection (see Breast Cancer Survival Component)  
Step 9: Generate the ages undergoing screening given birth cohort (see Screening Component)  
Step 10: Generate the tumor size detection threshold of mammography (see Screening Component)  
Step 11: Compute age, tumor size and stage at screen detection (see Screening Component)  
Step 12: Generate type of adjuvant therapy (see Breast Cancer Treatment Component)  
Step 13: Generate age at breast cancer death following screen detection given age and stage at breast cancer death (see Breast Cancer Survival)  
Step 14: Compute age of death as the minimum between age of breast cancer death and age of other cause of death  
Repeat for next individual  
Repeat for next birth cohort |
clinically detected with invasive breast cancer in the absence of the screening and other-cause mortality. The breast cancer patient is assigned the age that her first primary invasive tumor clinically surfaces.

This component relies on a CISNET base case input commonly referred to as the “secular trend in breast cancer incidence,” which was estimated from the historic Connecticut Tumor Registry (CTR) and Surveillance, Epidemiology, and End Results (SEER) (5). The base case input estimates incidence rate of the first primary breast cancer (invasive and in situ) in the absence of screening for annual birth cohorts starting 1891 by single year of age, for ages 25–84 years. The base case incidence is assumed to be zero for women younger than 25 years and older than 84 years.

We treat the base case incidence as the hazard rate of the first cancer and apply it to our estimated cancer-free population. For each birth cohort we interpret the given incidence per 100,000 women ($h_0$) as the hazard rate for age $a$. To reduce the computational time for multiple runs, we generate and sample from a distribution function of the clinical detection age ($A_{BC}$) at symptomatic detection of the first invasive breast cancer for each birth cohort as follows:

$$P(A_{BC} \leq a) = 1 - \prod_{i=25}^{a} (1 - h_i/100000)$$

where the age $a$ is an integer. Because this is a discrete distribution function of the woman’s age at the first symptomatic detection, we generate the exact age by assuming a uniform distribution within a year. The same calculation is made for all the birth cohorts.

Because the base case input provides the estimated trend for the sum of clinically detected in situ and invasive cases, we modified it to estimate the incidence for clinically detected invasive disease only. We adjusted it by removing an estimated proportion of ductal carcinoma in situ (DCIS) cases as a function of age from Surveillance, Epidemiology, and End Results (SEER) 1975–1979 data. The same correction factor was applied to all birth cohorts. It is possible that temporal and/or birth cohort variations affect this relationship, but we do not have informative data to produce such estimates. If we are overestimating the proportion of DCIS, we are overestimating the incidence and mortality of invasive breast cancer, and vice versa.

Further, the CISNET base case input on breast cancer secular trend exist may be overestimating the true hazard of the first primary, particularly in the older age groups. The base case incidence was approximated as the observed count of “new cancers” divided by the size of the midyear population based on data from the CTR and SEER, whereas the true incidence is defined as the count of “first cancers” divided by the size of the cancer-free population (6). This approximation is made because SEER does not include the size of the cancer-free population. It would produce the true incidence if women are equally at risk for breast cancer regardless of their history of breast cancer. However, the risk of breast cancer probably increases with a prior history of primaries as evidenced by the Gail model (7); hence, the approximation biases toward a larger number of first primaries and a possibly a larger breast cancer mortality. Its exact effect on breast cancer mortality is not obvious without taking into account the differences in the stage distribution and survival of a first primary and subsequent primaries.

### Natural History Component

Our natural history component specifies the size of a breast cancer patient’s first primary invasive breast tumor and its SEER historic stage from the moment the tumor is invasive and 2 mm in diameter to the moment it clinically surfaces. It also specifies the mechanism under which the first primary invasive tumor clinically surfaces. (By “clinically surfaces” we refer to the event due to a patient experiencing physical symptoms associated with breast cancer.) In this section, we describe the parameterization of the natural history model and parameter estimation.

#### Tumor size and stage during preclinical period

The tumor is assumed to be spherical and grow exponentially during its preclinical period. We define the preclinical period from the moment the tumor is invasive and 2 mm in diameter to the moment it clinically surfaces. The volume of the tumor at time $t$ is expressed as $V(t) = c_0 \exp(t/R)$, where the inverse growth rate $R$ [which is the doubling time divided by $\ln(2)$] has gamma distribution with rate $\alpha$ and shape $\beta$, as suggested by others (8).

We assumed that the disease starts in the local stage and progresses to regional and distant stages as the primary tumor increases in size (9). We define the onset of the regional stage as the point at which nodal involvement first becomes detectable by methods commonly used in clinical practice. Similarly, we define the onset of the distant stage as the point at which distant disease first becomes detectable by techniques commonly used in clinical practice. If the tumor is clinically detected before the onset of the regional or distant disease, it is staged as local disease. If the tumor is clinically detected after regional transition but before distant transition, it is staged as regional disease. If the tumor is clinically detected after the distant transition, it is staged as distant disease.

The hazard of the time to onset of observable regional disease is analytically modeled as

$$P(T_N \in [t, t+dt) | T_N \geq t) = (\eta_0 + \eta_1 V(t)) dt + o(dt)$$

This event is modeled as two processes: the hazard of the first process is constant over time; the hazard of the second process is proportional to the volume of the tumor at time $t$, $V(t)$ and therefore increases as the tumor grows in volume. Similarly, the hazard of the time to onset of observable distant metastasis ($T_M$), measured from the onset of the observable regional disease, is

$$P(T_M \in [t, t+dt) | T_M \geq t, T_N = t_N) = \begin{cases} (\omega_0 + \omega_1 V(t)) dt + o(dt), & t \geq t_N \\ 0, & t < t_N \end{cases}$$

We do not include a temporal trend in the onset of regional and distant disease due to the lack of informative data; yet, it is possible that technology advancements have caused a stage migration (10).

#### Clinical detection function

The hazard of the time to clinical detection ($T_D$) is assumed to be proportional to the tumor volume,

$$P(T_D \in [t, t+dt) | T_D \geq t) = \gamma V(t) dt + o(dt)$$

This clinical detection function was considered previously for breast cancer (8).

### Estimation of parameters in natural history model

Our natural history model has seven parameters: $\alpha$, $\beta$, $\eta_0$, $\eta_1$, $\omega_0$, $\omega_1$ and $\gamma$. Maximum likelihood estimates for these parameters are based on SEER data of the tumor size and stage of invasive...
cancers that were clinically detected in the absence of screening. Only SEER breast cancer cases detected between 1975 and 1981, which represents a period of no to little screening, were considered, and of these only the first primary tumor in a woman with multiple primaries was selected. Because we do not directly incorporate data that contain temporal information (such as the patient’s age), the rate parameter estimates are dimensionless. We scale them by the mean tumor volume doubling time (9). Estimates under the condition $\alpha = \beta$ [or similarly, a mean tumor volume doubling time of ln(2) years] are presented in Table 2, stratified by age groups (20–39 years old, 40–49 years old, 50–69 years old, 70–84 years old).

The mean tumor volume doubling time is estimated simultaneously with the median tumor size detection function, as discussed later.

**Assumptions about DCIS.** DCIS is not included in our natural history model because there is little known about its progression. Some forms of in situ (in particular, high-grade DCIS) have been suggested to progress to invasive disease (11), but what proportion progresses and how quickly it progresses are not known. For this reason, our model is limited to disease that would have been clinically detected as invasive. By not including clinically detected in situ disease, we are implicitly assuming that DCIS would have been clinically detected as invasive but is screen detected as in-situ. In our model, such cases of screen-detected DCIS are most likely captured as localized, small invasive tumor and as such would have a similar prognostic outcome to screen detected DCIS.

**Screening Mammography Component**

Our screening component specifies the screening dissemination and screen detection. With the screening component added, we have a means to estimate the mean tumor growth rate that was specified in our natural history model earlier.

**Screening dissemination module.** We use the CISNET base case input for mammography dissemination, which describes an increasing uptake of screening mammography for women aged 40 years and older. This input generates the ages at which an individual undergoes screening, given her birth cohort (12). We truncate the screening schedule at the age of clinical detection or death of other causes, whichever occurs first.

**Screen detection mechanism.** Each simulated woman who receives at least one screening examination is assigned a randomly generated mammographic detection threshold. The mammography detection threshold is defined as smallest tumor diameter detectable on screening mammography. Tumors below this diameter are missed and tumors above are classified as screen detected if they have not clinically surfaced before the time of the screening examination. Because the tumor size increases between screening examinations, the probability of screen detection increases.

Once a patient is screen detected, her age, tumor size, and SEER historic stage at detection are recorded. A patient is classified as an “interval case” if her tumor is clinically detected between two scheduled screening examinations, as illustrated in Fig. 2, B.

The distribution for the mammography threshold was modeled by assuming that the hazard function for “screen detectability”, i.e., the transition from a non–screen detectable tumor to a screen-detectable tumor, is proportional to the cross-sectional area of the tumor, which is in turn proportional to the tumor volume raised to the two-thirds power, i.e.,

$$P(V_{TH} \in [v, v+dv] | V_{TH} \geq v) = \lambda v^{2/3} dv + o(dv)$$

In terms of the tumor diameter, the resulting cdf is $F_{TH}(d) = 1 - \exp(-0.6(\pi/6)^{1/3}d^3)$. In our simulations, the distribution was truncated at diameter $d = 2$ mm, so that a tumor with diameter less than 2 mm would not be screen detected.

Our mammography detection function has the advantage that it is fully specified by one unobservable parameter but has the disadvantage that it produces a narrow distribution for the tumor size detection threshold. A wider distribution is more plausible; however, it would require an additional unobservable parameter that could not be identified from the available data.

**Joint estimation of median detection threshold and the mean growth rate.** We estimate two unobservable parameters, namely, the median tumor size detection threshold of mammography and mean growth rate simultaneously, by calibrating to the SEER incidence trends and data from the Breast Cancer Surveillance Consortium (BCSC), using a two-step procedure. In the first step, each of 5-year age-specific SEER incidence curves are smoothed with respect to the year of diagnosis by using natural splines (PLUS 6.1) in terms of the number of new cancers divided by the midyear population. Using our simulation program, we estimated incidence as a number of first cancers divided by the midyear population minus the prevalence. Sum of squared differences between age-specific smoothed SEER incidence and simulated incidence was used as a goodness-of-fit measure, assuming the same weight for each age group and each calendar year. This measure was computed over the two-dimensional parameter grid with increments of 0.05 year for the mean doubling time and 0.05 cm for the median threshold. The mean doubling time was varied between 0.2 year and 1.1 year and median detection threshold was varied between 6 mm to 12 mm. Various combinations of the parameters produced similar goodness-of-fit measures. In the second step, for each fixed threshold we selected the best mean doubling time. Using thus created “pairs,” we selected the one that fits better the median size at detection for screen-detected cases in BCSC 1994–2000 data of cancers screen detected within 3 years of the previous screening mammogram for women aged 50–69 years. The resulting estimates currently used are 0.75 year for the mean tumor volume doubling time and 1.0 cm for the median threshold of screening mammography.

**Adjuvant Treatment Component**

Our adjuvant treatment component identifies the use of adjuvant treatment and the corresponding survival benefit.

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**Table 2. Logarithms of the natural history model parameters, stratified by age groups**

<table>
<thead>
<tr>
<th>Age group, y</th>
<th>Log(α = β)</th>
<th>Log(γ)</th>
<th>Log(ηα)</th>
<th>Log(ηγ)</th>
</tr>
</thead>
<tbody>
<tr>
<td>20–39</td>
<td>−0.0974</td>
<td>−9.82</td>
<td>−10.11</td>
<td>−6.12</td>
</tr>
<tr>
<td>40–49</td>
<td>−0.129</td>
<td>−9.71</td>
<td>−9.96</td>
<td>−5.98</td>
</tr>
<tr>
<td>50–69</td>
<td>−0.0226</td>
<td>−9.74</td>
<td>−9.91</td>
<td>−4.69</td>
</tr>
<tr>
<td>70–84</td>
<td>0.0777</td>
<td>−9.82</td>
<td>−10.13</td>
<td>−4.31</td>
</tr>
</tbody>
</table>

*Estimates obtained for $\alpha = \beta$ or, equivalently, a mean tumor volume doubling time of ln(2) years.
**Treatment dissemination.** Each breast cancer patient is assigned adjuvant treatment (tamoxifen, multiagent chemotherapy, both, neither) depending on the patient’s age, tumor size, stage, estrogen receptor (ER) status, and year at detection, as specified by the NCI treatment dissemination base case input (13).

Tamoxifen dissemination targets ER-positive tumors in more recent years. Because ER status was not part of the natural history model, we assume that ER status does not vary over the preclinical course of the disease and does not affect the probability of screen detection. The probability that an individual breast cancer patient has an ER-positive tumor was based on the proportion of women with ER-positive disease in the SEER data 1990–1994: The proportion is 62%, 75%, and 83% for women younger than 50, 50–69, and 70 years or older at detection, respectively.

**Treatment efficacy.** We assume proportional benefits due to adjuvant treatment using published hazard ratios (14–16). For chemotherapy, the hazard ratio for the breast cancer–specific survival depends on the age at detection: 0.72 for women aged less than 40 years, 0.73 for women aged 40–49 years, 0.84 for women aged 50–59 years, and 0.91 for women aged 60 years and older. For tamoxifen, the benefit is assigned only to patients with ER-positive tumor and depends on the number of years that the treatment is assigned (2 years versus 5 years). The associated hazard ratios used are 2 years of tamoxifen and 0.69 for 5 years of tamoxifen. These hazard ratios were modified by 10% to convert the reported overall survival effect due to tamoxifen to a breast cancer–specific survival effect. If a woman receives both chemotherapy and tamoxifen, the product of hazard ratios is applied.

**Survival/Mortality Component**

Our survival component specifies a patient’s breast cancer survival time from the moment her first primary invasive tumor is detected. Breast cancer survival depends on the patient’s age, tumor size, and SEER historic stage at detection, as well as the mode of detection and use of adjuvant therapy.

**Baseline breast cancer survival.** Breast cancer survival curves for patients detected in the absence of screening and adjuvant therapy are a CISNET base case input, referred to as baseline breast cancer survival curves. These are Kaplan–Meier estimates obtained from SEER data for female breast cancer patients who were detected between 1975 and 1979 and for whom breast cancer was their first primary tumor. Because 1975–1979 is associated with minimal levels of screening and adjuvant therapy, the baseline survival curves are assumed to capture only the effects of primary breast cancer treatment, namely, surgery with the possibility of radiation. The curves are stratified by age at detection (i.e., 30–39 years old, 40–49 years old, 50–59 years old, 60–69 years old, 70–84 years old) and SEER historic stage (i.e., local, regional, distant); local and regional stages curves are further stratified by tumor size (i.e., <2 cm, 2–5 cm, and ≥5 cm). The tail of the Kaplan–Meier curve is assumed to represent a cure fraction.

**Breast cancer survival post–screen detection without adjuvant therapy.** The breast cancer survival curve post–screen detection is taken to be the maximum of two curves: 1) the baseline survival curve that corresponds to the age, size, and stage at screen detection and 2) the baseline survival curve that corresponds to the age, size, and stage at clinical detection. Both survival curves are initiated at the corresponding age of detection, and the probability of survival for clinical detection is set at 100% during the lead time. This approach rules out the possibility of death during the lead time.

The assignment of breast cancer survival post–screen detection is arbitrary. Lower breast cancer mortality rates would be obtained by using the baseline breast cancer survival curve that corresponds to the screen-detected tumor characteristics initiated at the age of clinical detection. Higher breast cancer mortality rates would be obtained by using the baseline breast cancer survival curve that corresponds to the screen-detected tumor characteristics initiated at the age of screen detection, because it would allow for death in the lead time. In a sensitivity analysis, we found that these two extremes do not produce outcomes that deviate substantially from the decision rule that we applied.

**Breast cancer survival following adjuvant therapy with and without screening.** We assume a proportional hazard reduction in breast cancer mortality due to adjuvant treatment (see “Treatment efficacy”). In the absence of screening, the hazard ratio is applied to the base case baseline breast cancer curves. In the presence of screening, the hazard ratio is applied to the resulting breast cancer survival curve obtained in “Breast cancer survival post–screen detection without adjuvant therapy.”

**Model Verification**

Three approaches were taken to demonstrate that our simulation model produces known or expected outcomes. First, we verified that the base case input distributions were correctly reproduced by our simulation, in particular, the distribution of time to death from breast cancer not subject to death from other causes; the distribution of time to death from other causes; the age-specific breast cancer incidence by birth cohort; the distribution of the time to first mammogram; the proportion of women receiving 1, 2, 3, 4, 5, 6, and 7 or more mammograms from 1994 to 2000; and the treatment dissemination profile. We also verified that the model-specific input distributions were correctly reproduced by our simulation, in particular, the distribution of clinically detected tumor sizes, the tumor size at nodal and distant involvement, and the growth rate distribution.

Second, we ran a series of hypothetical, diagnostic tests and verified that our simulation produces reasonable responses. We considered the following scenarios: a breast cancer risk of zero, no screening, 100% annual screening for women aged 40–79 years, low median detection threshold of mammography (2 mm), large median detection threshold of mammography (5 cm); low mean tumor volume doubling time (1 month); high mean tumor volume doubling time (18 months), and a flat secular trend.

Third, we performed internal validation tests and verified that our simulation produces reasonable responses. We analyzed the growth rate and tumor size distributions of screen-detected and interval cases to confirm that screen-detected cancers are more likely to be smaller and slower growing than interval cancers.

**Model Validation**

We evaluate our simulation model based on how well it predicts the observed U.S. breast cancer mortality rates from 1975 to 2000.

**Comparison of Predicted and Observed U.S. Breast Cancer Mortality**

Figure 3, B compares the predicted and actual age-adjusted breast cancer mortality rates from 1975 to 2000. The general
Higher than the observed level because we may be overestimating absolute level of the predicted breast cancer mortality rate. The sensitivity analyses are larger among women aged more than 60 years at death. The mortality is higher than expected because the incidence for the first primary may be too high (see "Breast Cancer Incidence Component"). We adjust the number of new first cancer patients as follows: 1) we determine if a patient has been diagnosed previously given her age and stage, from data from the CTR from 1975–1979; 2) if yes, then we return her to the healthy population. Although this adjustment may underestimate incidence of first cancer, we find that it does not change the shape of the predicted mortality trend but rather brings the absolute level closer to the observed level, as shown in Fig. 4, A.

Adding a fraction of screen-detected invasive tumors to be reclassified as screen-detected DCIS. Because we did not account for the possibility that some invasive tumors could have been screen detected as DCIS, we may have overestimated breast cancer mortality. We suspected that the small change in mortality may be due to an improvement in the tumor size detection after the year 1995. Despite the significant improvement in baseline survival, we do not conclude that an improvement in tumor size detection function of mammography would not substantially affect our results.

Adding a temporal trend to baseline survival. The CISNET base case input for age- and stage-specific breast cancer survival following primary surgery did not include a temporal trend. To evaluate the impact of such a trend, we considered a stepwise change in the baseline survival by forcing a 20% improvement in a specific calendar year. The results from a stepwise change in years 1985, 1990, and 1995 are shown in Fig. 4, B. The change in mortality was immediate. Because even a gradual improvement in baseline survival could substantially alter the shape of the predicted mortality curve before the year 1995, we do not consider it to be the major cause for the discrepancy.

Adding a temporal trend to mammography detection. We did not incorporate a temporal variation in the detection function of screening mammography due to the lack of informative data. To evaluate the impact of such a trend, we introduced a stepwise change in the median tumor size detection threshold of mammography by reducing it from 1.0 cm to 0.5 cm at a specific calendar year. The results from a stepwise change in years 1985, 1990, and 1995 are shown in Fig. 4, C. A noticeable change in breast cancer mortality begins approximately 3 years after the stepwise change. Despite the significant improvement in the detection threshold, the reduction in mortality is not large enough to account for a mortality decline after the year 1995. We suspected that the small change in mortality may be due to binning the base case breast cancer survival curves, with the single, best prognostic curve associated with all localized tumors less than 2 cm. When we separated survival curves for localized tumors less than 1 cm and between 1 cm and 2 cm, we still did not find a large effect on breast cancer mortality. We conclude that an improvement in the tumor size detection function of mammography would not substantially affect our results.

Sensitivity Analysis

We performed sensitivity analyses to evaluate the influence of modeling assumptions on the absolute level and shape of the predicted U.S. breast cancer mortality rate.

Varying the secular trend in breast cancer incidence. The absolute level of the predicted breast cancer mortality rate is higher than the observed level because we may be overestimating the true hazard of first cancer, as discussed in “Breast Cancer Incidence Component.” We adjust the number of new first cancer patients as follows: 1) we determine if a patient has been diagnosed previously given her age and stage, from data from the CTR from 1975–1979; 2) if yes, then we return her to the healthy population. Although this adjustment may underestimate incidence of first cancer, we find that it does not change the shape of the predicted mortality trend but rather brings the absolute level closer to the observed level, as shown in Fig. 4, A.

Adding a temporal trend to treatment efficacy. The CISNET base case input for efficacy of adjuvant therapy did not include a temporal trend due to the lack of supporting data. However, a
variation in efficacy is plausible given increasing variety of chemotherapeutic regimens over time. To evaluate the impact of such a variation, the efficacy in adjuvant treatment was assigned a stepwise change by improving the efficacy by 2 standard deviations (SD) based on published meta-analysis (15, 16) in a specified calendar year. The results following a stepwise change in 1985, 1990, and 1995 are shown in Fig. 4, D. A decrease in breast cancer mortality was immediately observed after an improvement in treatment efficacy. Because the dissemination of adjuvant therapy increases over time, we found that a linear change in treatment efficacy, ranging from −2 SD in 1976 to +2 SD in 2000, improves the model’s fit in the later years without substantially affecting the fit in the earlier years, as shown in Fig. 4, E. The lack of temporal change in treatment efficacy could explain the unpredicted decline after 1995.

Discussion on our Model Fit to Observed U.S. Breast Cancer Mortality

Our model reproduces the shape of the breast cancer mortality rates from 1975 to 1995. The flattening of our predicted breast cancer mortality rate from 1976 to 2001 that does not agree with the observed decline. We suspect that the lack of temporal variation in many of the base case inputs is the major cause of this discrepancy. After performing sensitivity analyses, we concluded that temporal variation in adjuvant treatment efficacy is probably the major reason for the discrepancy between the modeled and observed mortality after 1995. However, it is possible that multiple temporal variations can be included simultaneously in a manner that improves the fit to breast cancer mortality.

Although it was tempting to add temporal variations to the base case inputs and estimate additional model parameters by calibrating to breast cancer mortality trends, we did not do so for several reasons. First, calibration to mortality would have compromised our only means for evaluating our model’s performance. Second, it is not clear which temporal variations to add and how to best parameterize them; too many parameters would overparameterize our model. Third, calibration to mortality would force dependences among the model components, thereby increasing the risk of propagating potential errors due to misspecification in individual model components. Fourth, calibration would have made our model less transparent, as changes to the inputs could not be easily mapped to changes in the output. Fifth, calibration to mortality would have forced us to hypothesize that our model explains all the factors underlying the mortality.

Fig. 4. A) Breast cancer mortality rate based on National Center for Health Statistics data (NCHS) (solid black curve), the simulation model (solid gray curve), and the simulation model with the incidence modified (dashed gray curve). B) Breast cancer mortality rate when a stepwise change in baseline breast cancer survival occurs in 1985 (curve A), 1990 (curve B), and 1995 (curve C). C) Breast cancer mortality rate when a stepwise change in the mammography tumor size detection threshold occurs 1985 (curve A), 1990 (curve B), and 1995 (curve C). D) Breast cancer mortality rate when a stepwise change in treatment efficacy occurs in 1985 (curve A), 1990 (curve B), and 1995 (curve C). E) Breast cancer mortality rate following a linear temporal change in treatment efficacy (curve A), from −2 standard deviations (SD) in 1976 to +2 SD in 2000. Breast cancer mortality rate based on NCHS data (solid curve) and the simulation model using base case inputs (solid gray curve) are included in each figure. All rates are age adjusted to the 2000 U.S. standard population.
trends. Finally, calibration would have forced us to modify the CISNET base case inputs and thereby make our model not directly comparable to other CISNET models.

Even though we did not include DCIS in our base case analysis, we produced a reasonable match to breast cancer mortality. If screen-detected DCIS has had a dominant impact on breast cancer mortality, then we should have generated a much worse prediction of breast cancer mortality since DCIS accounts for a substantial fraction of incident breast cancer in the presence of screening. Breast cancer that would have been clinically detected as invasive disease in the absence of screening but is detected as DCIS in the presence of screening is probably being represented in our model as small, localized tumors with comparable prognosis. However, this finding does not imply that screen detecting disease as DCIS instead of small, localized disease is not clinically relevant. A more in-depth analysis needs to consider the complex detection characteristics of mammography for in situ versus invasive cancer.

CISNET Base Case Result

We now turn our attention to the CISNET base case question: “What are the contributions of screening and treatment to mortality reduction?” Fig. 5 illustrates our predicted age-adjusted breast cancer mortality trends from 1975 to 2000 under the following four scenarios: 1) in the absence of screening and adjuvant therapy, 2) in the presence of screening only, 3) in the presence of adjuvant therapy only, and 4) in the presence of both screening and adjuvant therapy. In the absence of screening and treatment, we predict a steady increase in age-adjusted breast cancer mortality due to the secular trend in incidence. Compared with the predicted mortality rate in the absence of screening and adjuvant therapy in the year 2000, the mortality rate in the presence of both screening and adjuvant therapy is reduced by a total of 29.9%, which is broken down as follows: 16.9% due to screening, 6.9% due to chemotherapy, and 8.9% due to adjuvant therapy. The estimated relative contributions of screening and adjuvant therapy to the mortality reduction were similar in magnitude: 53% due to screening versus 47% due to adjuvant therapy. A closer match to the absolute level of the breast cancer mortality rate would not affect these results. However, because our model does not match the declining rate of breast cancer mortality after 1995, we may be underestimating the effect of screening and treatment. On the basis of sensitivity analyses, we find that our result is robust for the benefit of screening mammography but may be underestimating the benefit of adjuvant therapy.

Conclusion

Our simulation model predicts the shape of the U.S. breast cancer mortality rates from 1975 to 1995 closely but predicts a leveling of the rates after 1995 instead of a decline as observed. Our model also predicts a slightly higher breast cancer mortality rate than observed. Through sensitivity analysis, we found that the higher predicted mortality rate may be the result of overestimating the rate of first cancer. The leveling off of breast cancer mortality predicted after 1995 is most likely due to a lack of temporal variations in many of the model inputs. However, without informative data, it is not clear which among many possible modifications to our model would be most appropriate to yield a closer fit to the observed breast cancer mortality trends. The discrepancy between the predicted and observed mortality rates can be considered an estimate of the magnitude of the effect from factors that were not explicitly modeled.

We predict that the contribution of screening and adjuvant therapy were roughly equal in magnitude. Our findings are independent of the screening mammography trial data and thereby provide alternative evidence of the mortality reduction due to screening. Because our predicted mortality after 1995 is greater than observed, we may be underestimating the benefit of screening and adjuvant therapy. Through sensitivity analysis, we found that our results for screening are robust and that the unpredicted mortality decline after 1995 is most likely due to temporal trends in the efficacy of adjuvant therapy that were not included in the base case input due to data limitations. Our model can be easily updated as new data becomes available to better inform the inputs.

References


NOTE

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