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Presumed benefit: lessons from the American experience with marrow transplantation for breast cancer

H Gilbert Welch, Juliana Mogielnicki

Few stories in medicine are as sobering as the American experience with autologous bone marrow transplantation (ABMT) for treating breast cancer. It is a story of young women dying from aggressive disease, well meaning physicians trying to be equally aggressive in treating it, and lawyers arguing that insurers should pay the bill. It is also a story of professional interests, weak research, financial gain, politics, and fraud. Because of its potential relevance to complex cancer therapies currently in development (such as gene therapy) we recount here the story and its lessons.

Early reports
Bone marrow transplantation was first performed to treat primary bone marrow disorders, but in the late 1970s it started to be used also for “rescuing” patients (using their own marrow) after supralethal chemotherapy or radiation for solid tumours.1 By the mid-1980s there were strong proponents for using it this way in advanced breast cancer—on the basis that higher chemotherapy doses would be expected to kill more tumour cells. The enthusiasm for this hypothesis was evident in comments made to the New York Times in 1989 by the head of the breast cancer section at the National Cancer Institute: “The evidence is absolutely convincing that the dose intensity is correlated with survival.”2 But other oncologists were more sceptical. In the same article one pointed out that the notion of dose-response was purely theoretical and also applied to toxicity: “It’s a hypothesis . . . and higher dosages are more toxic.”

And there were few data. The first report in the general medical literature on the treatment’s efficacy for breast cancer appeared in the Annals of Internal Medicine in 1988.3 The article reported on 172 women from 27 studies. The summary response rate (defined as tumour shrinkage ≥50%) was 58%. There were no controls. A few months later the Annals published a review that included 159 women and noted an 80% response rate.1 In both articles the authors were cautious, concluding: “Critical evaluation will require controlled trials” and “response rates that are probably superior to the best available with conventional therapies . . . although not yet associated with improved survival.”

Comments made to the press, however, were less cautious. In the news section of the Journal of the National Cancer Institute, one author said: “I think this shows that ABMT can be a very effective form of treat-

Summary points
For over 10 years bone marrow transplantation for breast cancer was seen as an example of the general dilemma about who should pay for costly new life saving therapies

This characterisation obscured the more basic question: Did it work?

Intermediate outcomes and inadequate controls made preliminary evidence misleading

Statements by physicians in the literature and the general press reinforced the presumption of benefit, as did the decision of government bodies to mandate insurance coverage

The findings of major randomised trials did not support the use of the therapy

This experience provides lessons relevant to complex cancer therapies currently in development

Patients in the media
Other factors converged to make bone marrow transplantation for breast cancer a big story. Women’s issues were prominent. Breast cancer was both common and feared. Transplantation was a source of hope—a technologically advanced procedure. And soon there was the added dimension of money. In December 1989 the Washington Post described a Johns Hopkins study of 20 women treated with bone marrow transplantation and reported “partial success” (while acknowledging that the prospects for long term survival were not yet known).4 The article reassured readers that the high cost of the treatment ($75 000-100 000) “is usually covered by health insurance or Medicaid.”
Four months later a 35 year old mother of three proved this statement wrong. Page 1 of the Post's Metro section declared: “Maryland Mother's Chance of Life Hinges on Trial; Patient Sues Insurers For Cancer Treatment Cost.” Pamela Pirozzi had been advised that her “best chance of surviving more than a year” was a transplant, but her insurance company had refused coverage, stating the procedure was still “experimental.” Armed with a “list of insurance companies in other states that, when challenged, have agreed to pay for the procedure,” the Pirozzis sued.

A federal judge ruled in her favour: “To require that the plaintiff or other plan members wait until somebody chooses to present statistical proof . . . that would satisfy all the experts means that plan members would be doomed to receive medical procedures that are not state of the art.” The same month another federal judge ordered a Massachusetts insurer to pay for a Boston woman to receive a transplant in North Carolina.

State of the art or state of the theory?

American insurance companies argued that bone marrow transplantation was experimental and thus not covered by their policies: “We view ABMT for breast cancer as investigational and experimental, because the treatment has not proved to be safe and effective.”

Such a broad definition did not fare well in courts or the court of public opinion. It also highlighted the dilemma for policy makers: how do you define “experimental therapy”? In November 1990 the Blue Cross Blue Shield Association of insurers provided a pragmatic definition: any therapy being studied in a randomised trial.

After years of inconclusive data a National Cancer Institute panel had just decided to conduct a major randomised trial to test the effectiveness of bone marrow transplantation. The national study was to recruit 1200 women with metastatic breast cancer or high risk regional disease. The association promised at least $10m to help fund the trial.

It was the first time a private insurer had agreed to fund a trial of treatment. The decision was a reaction to the controversy and the lawsuits. The senior vice president of the association pointed out the benefits of the decision: “By funding some of the clinical care costs associated with the National Cancer Institute trials, we're able to provide access to this treatment while awaiting trial results, the presumption of benefit was also clear: “Using reasonable assumptions, ABMT provided substantial benefit but at a cost that may be untenable.”

Health services research and ABMT

There was sufficient controversy in the medical community to attract non-oncologists. In April 1992 the Journal of Clinical Oncology published a structured review of the data by an epidemiologist: 72 studies and abstracts on the use of bone marrow transplantation for breast cancer—but no randomised trial. After applying three simple inclusion criteria (peer review, survival or response as an outcome, and more than 10 patients), the reviewer had only 10 case series left to summarise (238 patients). Six large series of conventional therapy and 45 randomised trials (comparing various agents) served as a crude comparison group. The author concluded that survival rates after transplantation and conventional chemotherapy were essentially the same.

The same month JAMA published an article on bone marrow transplantation for breast cancer—ironically addressing the cost effectiveness of the procedure. Given that there were no good data on effectiveness, this was predicted using case series data on response rates and a decision analytic model. In retrospect the model's predictions were half right. The estimated 27% three year survival after transplantation was similar to that observed in the subsequent randomised trial, but the estimate of 14% survival for those treated with conventional chemotherapy was less than half that actually observed. The reported cost effectiveness ratio was $115 800 per life year. Although the authors were clear that these data were to be used while awaiting trial results, the presumption of benefit was also clear: “Using reasonable assumptions, ABMT provided substantial benefit but at a cost that may be untenable.”

The juxtaposition of these two articles was striking. On the one hand was the lack of evidence of effectiveness. On the other was the conclusion that it was effective but too expensive. The authors of the two articles tried to reconcile the “apparently different conclusions” in a letter to JAMA, but the sound bite was already out: “High cost marrow treatment helps fight breast cancer.”

Insurance coverage and lawsuits

Despite the controversy about whether the therapy worked, more women were taking their insurers to
Politics and policy

The presumption of benefit was widespread. Because investigators were struggling to enrol patients in the randomised trials, even the mundane issue of patient accrual made headlines.26 Four years into enrollment, the trial of patients with metastatic breast cancer had only half of the 549 patients needed. Many physicians and patient advocacy groups believed the question was not in doubt: transplantation was better.27 Others pointed out the strong financial incentives for medical centres to perform transplants.28

In September 1994, while the National Cancer Institute was arguing that there was scientific uncertainty about the effectiveness of transplantation, a different branch of the government weighed in.29 The Office of Personnel Management ordered all health plans serving federal employees to grant coverage of autologous bone marrow transplantation for breast cancer within 24 hours or risk being dropped from the programme. The decision affected 350 health plans serving 9 million people.

The decision was officially based on several factors. Nearly a third of the programme's health plans already covered the procedure or were about to. Virginia, the home of many federal employees, had recently mandated state-wide coverage (one of seven states ultimately to do so). Finally, the increasing number of lawsuits highlighted the downsides of not covering the procedure.

There was also political pressure. In October 1993 54 members of congress wrote to demand that the Office of Personnel Management cover the procedure.30 They cited statistics from a report at Duke University claiming transplantation and high dose chemotherapy were "eight times more effective than conventional dose therapy." (The statement was loosely based on an article on bone marrow transplantation in high risk primary breast cancer reporting a 72% 5-year survival rate compared to 35% in historical controls.)23 In June 1994 the directors of five major academic cancer treatment programmes wrote to the office, presenting even more favourable data. Finally, Representative Eleanor Holmes Norton led a hearing in August 1994, with testimonies from federal employees who had been denied coverage. Despite hearing the National Cancer Institute position that "formal scientific evaluation (ought) to proceed the routine use of such a toxic procedure."31
and expensive therapy she called for the Office of Personnel Management to re-evaluate its policy.

**Fraud and findings**

In 1995 the *Journal of Clinical Oncology* reported on the first randomised trial of bone marrow transplantation in metastatic breast cancer. South African oncologists led by W R Bezwooda reported a complete response rate (no evidence of tumour) of 51% in women randomised to transplantation, compared with 4% in those receiving conventional therapy. The benefit in median survival was even more impressive: 90 weeks versus 43. The article was cited about 300 times before concerns about its validity. Four years later at the 1999 American Society of Clinical Oncology meeting, Bezwooda reported equally impressive results for transplantation in high risk primary breast cancer. At the same meeting, however, four other randomised trials were presented (Lotz JP; Peters W; Scandinavian group 525, >10 nodes 36 78 80; French group 874, stage IV 36 78 80; Philadelphia group 15 199, stage IV 36 78 80; Peters B, stage IV 36 78 80).

The obvious lesson from these events was articulated in the *New York Times* by two of the treatment's most visible critics. “As a society we have to accept that rigorous evaluation of a new treatment is essential … Skipping this step may seem like a compassionate act, but it can have devastating consequences.”

It is important to remember that preliminary evidence can be misleading; intermediate outcomes (such as response rates) may not correlate with survival, and historical controls may not be comparable. And proponents can be persuasive. The lesson is familiar: it is the case for randomisation. Future research may still show the utility of a chemotherapeutic regime for breast cancer which requires autologous bone marrow transplantation, but for now this story serves as a good example of why scepticism is important in medicine. There are also less obvious lessons.

**Lessons**

For over 10 years desperately ill women had sought bone marrow transplantation as their best chance for survival. Many physicians encouraged this judgment. Fearing bad publicity and lawsuits insurers reluctantly agreed to pay the considerable charges. A strong presumption of benefit and equally strong financial interests impeded progress towards finding an answer.

The media was slow to see that there was more to the story than the question of how to pay for a
costly, new, life saving therapy. Proponents were successful in characterising the case against transplantation as simply about money. Yet proponents of “advances” will always be more vociferous than detractors; they usually have stronger interests (both professional and financial) in arguing for a particular technology than detractors have in arguing against it. Given a public primed to believe in medical breakthroughs, the press should focus on evidence of effectiveness before raising arguments about money. All would be well served by a press that displayed the same scepticism about pronouncements from medicine as it does with pronouncements from government.

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Why does NICE not recommend laparoscopic herniorrhaphy?

Roger W Motson

NICE’s reluctance to recommend laparoscopic repair of hernias is based mainly on economic considerations, some of which are inaccurate, according to Roger Motson

More than 100 000 inguinal herniorrhaphees are carried out each year in the United Kingdom, making it one of the commonest operations. Newer techniques have superseded the simple suture technique popularised by Bassini more than 100 years ago: firstly the tension-free darn with monofilament nylon and then the Lichtenstein repair with a polypropylene mesh patch. Although there was no initial randomised trial of the Lichtenstein technique, it rapidly gained popularity during the past decade. Laparoscopic repair, which places a considerably larger polypropylene mesh patch against the inner surface of the abdominal wall than that used in the Lichtenstein technique, was first performed about 10 years ago. This larger patch reinforces the entire groin, covering the sites of both indirect and direct inguinal hernias and also of femoral hernias. The position of the mesh is the same as that used in the

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**Summary points**

Inguinal herniorrhaphy is one of the commonest operations in the United Kingdom

Laparoscopic herniorrhaphy is less painful postoperatively than traditional open repair and allows the patient to return to work more quickly

The true costs of laparoscopic repair are lower than those of open repair, particularly when it allows detection and simultaneous repair of an undiagnosed contralateral hernia

Surgeons are under-represented on NICE’s appraisal panel

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