Voices of a Broken System: Real People, Real Problems

President’s Cancer Panel
Report of the Chairman
2000–2001

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This report is submitted to the President of the United States in fulfillment of the obligations of the President’s Cancer Panel to appraise the National Cancer Program as established in accordance with the National Cancer Act of 1971 (P.L. 92-218), the Health Research Extension Act of 1987 (P.L. 99-158), the National Institutes of Health Revitalization Act of 1993 (P.L. 103-43), and Title V, Part A, Public Health Service Act (42 U.S.C. 281 et seq).

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Dear Mr. President:

Few families in America have not been touched by cancer. Yet a great many people—both the privileged and the poor—find that at the very time they need the most effective cancer care our research enterprise has devised, the health care delivery system of our Nation fails them.

People with cancer need your help now.

Mr. President, profound advances are being made in cancer research, but our health care delivery system is broken. It is the root of vast and unnecessary suffering, personal financial ruin, and loss of dignity for millions of people with cancer, who must fight their way into and through a dysfunctional system even as they struggle to save their very lives. The overwhelming majority of these people have worked and contributed to the nation throughout their adulthood. Many pay sizeable health insurance premiums every month, believing they are protecting themselves and their families, but may find themselves faced with unreimbursed cancer care costs that quickly drain their life savings. Many work hard every day, but cannot afford even basic health insurance. Some are unable to work, but are no less deserving of proper care when faced with a life-threatening illness.

In 2000 and 2001, acting according to its legislative mandate and in your behalf, the President’s Cancer Panel traveled to every region of the country to listen to the voices of cancer patients and survivors, family care givers, cancer patient advocates and volunteers, and to health care professionals who provide cancer care. The attached report summarizes their moving and unsparingly candid testimony, and presents the Panel’s recommendations for addressing the cancer care problems faced each day in every neighborhood in America.

These problems are many. They include lack of health insurance—a dire problem faced by 44 million people—inability to pay the out-of-pocket costs of cancer care, particularly oral medication costs; public and private health plan restrictions; physical distance from sources of care; and lack of transportation. Relatively few people with cancer receive full, accurate, and understandable information about their disease, either from their health care providers or from other sources. These problems stem primarily from insufficient provider communication or knowledge, and from language, literacy, and cultural barriers. Moreover, bias based on cultural and racial differences far too often causes some providers to offer less than optimal care, and causes some patients to avoid accessing the care they need because of fear or mistrust.

I hope that through the attached report, you too will hear these voices from every corner of America and take action—their voices could as easily be our own. The President’s Cancer Panel enjoins you to use the power and compassion of your office to direct all necessary Federal agencies to work together to implement the actions recommended in this report. I would welcome and look forward to an opportunity to meet with you to discuss this most urgent issue for the American public.

Sincerely,

Harold P. Freeman, M.D.
Chairman
Table of Contents

Executive Summary ................................................................. i

Preface .................................................................................... xv

Our Broken Health Care System Is Failing People With Cancer .................. 1
• Who Has Cancer in America?
• The Disconnect Between Cancer Research and Cancer Care Delivery
• The Health Care System and Cancer—Who Is Underserved?

What Is Happening to Real People: Findings ........................................ 7
• What Barriers Limit or Prevent Access to Care?
• What Barriers to Cancer Care Are Related to Information and Education?
• The Impact of Culture and Bias on Cancer Care

What Can Be Done to Help People Now: Recommendations .................... 37

References .................................................................................. 39

Appendices ................................................................................... 43
A. 2000–2001 President’s Cancer Panel Regional Meeting Locations and Participating States and Territories
B. Bringing Cancer Care Advances to the Public: Bridging the Divide Between Research and Delivery, Figure A

Appendices available online at: http://deainfo.nci.nih.gov/ADVISORY/pcp/minsmenu.htm
Minutes of 2000-2001 Meetings
The issues of cancer care in America are everyone’s issues. Each day, 34,000 people in America are diagnosed with cancer and another 1,500 die from the disease. Nearly nine million more are living with a cancer history. And every person, regardless of income, education, ethnicity, race, age, or geographic locale, is at risk of developing cancer.

Growing evidence indicates that most people in America receive neither the most appropriate care when faced with a cancer diagnosis, nor adequate cancer prevention and detection services. Factors contributing to this massive failing are many and complex, but the clear and central issue is the failure of our health care system to deliver, in an equitable and timely manner, the cancer care we know is most effective—regardless of a person’s insurance status or ability to pay.

The President’s Cancer Panel, established by the National Cancer Act of 1971 to monitor implementation of the National Cancer Program, previously reported to the President on this critical “disconnect” between our cancer research discoveries and the type, timeliness, and continuity of cancer care that people actually receive. In 2000 and 2001, the Panel held seven regional meetings to hear firsthand from people with cancer, their families, and the health professionals, administrators, advocates, and volunteers who serve them—393 in all—about problems they experience in accessing and providing cancer care and cancer information. In every corner of the Nation, patients and professionals alike echoed the same moral tenet:

No person in America with cancer should go untreated, experience insurance-related diagnosis or treatment delays that jeopardize survival, or be bankrupted by a cancer diagnosis.

Yet these very things are happening to far too many of us. The problems of cancer care in America are not theoretical, analytic, or abstract—they are real problems affecting real people.
What Is Happening to Real People: Findings

The numerous issues described by meeting participants fall broadly into two categories: access and information. In addition, cancer care can be influenced greatly by behaviors and decision-making by both health care providers and the public that stems from perceived differences among populations and individuals.

Barriers Limiting or Preventing Access to Cancer Care

Access barriers include those related to the organization and operation of the health care system itself, financial barriers to care, and physical barriers that reduce or prevent access. However, these categories are not always mutually exclusive.

System Barriers

The current health care system underemphasizes cancer prevention and often allocates cancer funding by disease site. Both approaches are counterproductive to providing comprehensive cancer care and developing effective cancer control programs. Cancer prevention, education, and screening efforts are limited at best and highly uneven across the country. Treatment for detected cancer remains unavailable to some of the uninsured unless they are able to obtain charity care or qualify for medical assistance. In addition, coordination between public payers is poor, and patients often are not informed of all health benefits for which they may be eligible. Believing they have no coverage or limited coverage, patients may incur unnecessary out-of-pocket costs, delay treatment, or even forego care.

System-related barriers to care most often described by people with employer-sponsored or other private health insurance include fragmentation of care, gatekeepers who control access to screening and specialists, and limitations or exclusions on specific drugs and services, including clinical trials. Numerous patients recounted having to fight their insurers to get the care they needed to save their lives. The current system also discourages appropriate end-of-life care, resulting in late referral to hospice (or no referral) and causing many terminal cancer patients to die without adequate pain and other symptom control.
Financial Barriers

Financial Issues Affecting Patient Access to Care. For people with cancer and their families, health care system issues that can be explained logically by fiscal and economic realities often translate into a personal reality that includes family bankruptcy, needless suffering, loss of dignity, and loss of life. Currently, 44 million Americans have no health insurance at all. Uninsured rates are as high as 25 percent of the non-elderly in some states, with much higher rates in some rural and frontier regions.

Many of the cancer survivors and family caregivers providing testimony were self-employed—farmers, ranchers, small business owners, and other independent workers. These speakers explained that they seldom can afford even basic health insurance, though they make enough money to support themselves and their families. The working poor may hold two or three jobs, none offering health benefits. This population is likely to lack both health insurance and the financial reserves to see them through an extended illness. They often avoid cancer screening or care for suspicious symptoms because they know they cannot pay for cancer care. Late stage diagnosis is a common result. When cancer strikes, uninsured workers may find they have too much income or too many assets to qualify for Medicaid or other medical assistance, but are too young for Medicare. These patients can quickly amass huge medical debts that will take the family many years to repay. Some are forced into bankruptcy.

To qualify for Medicaid, patients typically must give up the employment that provides family support and divest themselves of virtually all assets, including their car, home, business, or farm. In addition, survivors described such long delays in obtaining Medicaid approval that they were faced with more advanced disease by the time they were able to begin treatment. Even after securing a Medicaid card, it sometimes took weeks or months to find a provider who would accept Medicaid payment. Cancer patients who remain disabled by their disease for 12 months may qualify for Social Security Income (SSI) payments; however, this income may exceed Medicaid eligibility ceilings, causing them to lose access to cancer care.

At least 31 million non-elderly insured Americans are underinsured for cancer care costs. Though they pay substantial monthly premiums, many find the combined burden of copayments, deductibles, non-covered services,
medical supplies, and drug costs (particularly for oral chemotherapy and supportive medications), slow health plan reimbursements, and additional transportation and child care costs quickly exhausts family savings. This is particularly true when patient income is reduced or lost and/or when a family wage earner must work less in order to care for the cancer patient. Should the patient or spouse lose the job providing health insurance, the family can face a quick descent into indigent care and bankruptcy. Some survivors who return to work but have lapsed insurance coverage may find they are uninsurable, subject to a lengthy waiting period for cancer-related coverage, or eligible only for prohibitively expensive coverage.

Financial Issues Affecting Health Care Providers. As health care payers and purchasers struggle to contain health care costs, financial pressures on health care providers continue to increase, affecting the care available to people with cancer and those in need of screening, diagnostic, and preventive services. Survivors and family members reported widely varied experiences in terms of provider willingness to accommodate patients’ lack of insurance or full insurance coverage. Some patients were told to “pay what you can, when you can;” others were able to negotiate reduced rates and payment plans; and some providers donated their time and services. In other cases, however, providers asked for advance payments of $20,000 to $100,000 before initiating treatment. These highly divergent responses to patients in need appear to reflect an extreme and intensifying conflict between some providers’ commitment to render care as needed and a range of financial pressures that threaten the solvency, vitality, ethics, and integrity of health care institutions and individual providers alike.

Providers described ever-increasing payer-related paperwork that drives up administrative costs and reduces time available to see patients. Shrinking reimbursements and vastly different reimbursements for identical drugs and services in urban versus rural areas threaten the future of community oncology, particularly chemotherapy administered in office settings. These payment differences also are affecting rural providers’ capacity to upgrade equipment and staffing to provide more effective therapies in the community. As a radiation oncologist from the Midwest pointed out, “It costs the same amount of money for me to buy a linear accelerator as it does for somebody in New York City.” Highly disparate reimbursements are forcing ill and fatigued patients to travel long
distances to obtain care they could receive locally if provider reimbursements for that care were equitable in the rural setting. Lack of reimbursement for oral chemotherapy and supportive medications under Medicare and many private health plans also prevents patients from receiving care close to home. Speakers maintained that Medicare and Medicaid reimbursement cuts are increasing the gap between state-of-the-art cancer care and its implementation for all people. Testimony indicated that diminishing reimbursements also are a key factor driving increased pressure on physicians to see more patients each day. Physicians judged by local health plans to be inefficient may face financial penalties. While providers may suffer reduced revenues and greater stress, patients may suffer more costly losses: inadequate discussion of medical history and all relevant health issues; inadequate patient education; missed opportunities for preventive care; failure to be offered clinical trials or other care that requires additional explanation, monitoring, or paperwork; and in the worst cases, misdiagnoses.

Physical Barriers

Living in rural or frontier areas poses a special set of problems that comprise a vastly underappreciated barrier to cancer care access. Approximately one-fourth of the U.S. population lives in areas designated as rural or frontier, and half of all states have frontier regions (i.e., fewer than 6.6 people per square mile). Excessive distance from cancer care is due in part to the concentration of health care personnel and resources in urban areas, the lack of public transportation systems in rural and frontier areas, and the fact that many rural and frontier residents lack the resources to travel to care. The scarcity of both primary and specialty care providers in rural and remote areas is a longstanding problem that contributes to late diagnosis of cancer in these populations. Lower reimbursements for care provided in rural settings not related to operating cost differences and limited grant funding to sparsely populated areas continue to hamper efforts to recruit and retain oncologists and other cancer care professionals. Speakers called for incentives to attract health care professionals to these regions and help them sustain careers in underserved areas, including underserved inner cities.
Most rural areas have no public transportation system; patients reported driving up to 300 miles one way for screening or treatment, and some are too ill to do so. Because some services cannot easily be taken to rural areas (e.g., radiation oncology), patients may choose treatment options (e.g., mastectomy versus lumpectomy) that do not require repeated trips to a distant treatment facility. Native Americans, particularly those living on reservations, frequently drive long distances to an Indian Health Service (IHS) hospital or clinic and wait all day to be seen, only to be turned away and told to return the next day. This situation is so discouraging that some avoid seeking care for symptoms until they require emergency care. Many managed care enrollees and Veterans Administration beneficiaries must obtain services from contracted providers distant from the patient’s home, sometimes in another state.

Some patients lack reliable transportation or the money for fuel and tolls needed to travel to treatment or screening. Older Americans may not have a family member or friend able to drive them to medical appointments and may be unable to pay for transport. Transportation barriers are particularly onerous for patients in isolated areas such as Alaska, Hawaii, the Virgin Islands, Puerto Rico, and the Pacific Territories, where the only way to reach many, if not all, cancer screening or treatment services is by air or boat; these transportation costs may be prohibitive. In mountain states and rural northern regions of the country, reaching cancer care may become impossible for weeks at a time when roads become impassable due to winter weather. Such conditions can delay cancer diagnosis and disrupt treatment regimens. Telemedicine has the potential to bring some state-of-the-art cancer care services and continuing medical education to geographically isolated areas.

Transportation also is an issue for inner city residents who may not own cars or be able to afford bus or subway fares. In addition, some urban cancer patients are too sick to undertake a trip that may cover few miles but involves multiple bus transfers and considerable walking.

**Barriers Related to Information or Education**

Lack of information or education, not knowing how to find or evaluate information, not believing or acting appropriately on available information, not knowing how to get needed care within the health system—all can constitute barriers to cancer care.
Provider-Related Information and Education Barriers

Lack of information about cancer and cancer care was the provider-related barrier most commonly cited by those providing testimony at the regional meetings. Speakers emphasized strongly the need to better educate primary care providers about cancer. Initial decisions about care can be a critical determinant of patient outcome, and primary care providers, especially in rural and remote areas, often make these crucial referral and treatment choices.

Considerable disagreement exists, even among oncology professionals, about sometimes conflicting published screening and treatment guidelines. More broadly, there is confusion among providers, payers, and patients as to what constitutes quality care for cancer.

Speakers graphically described the serious repercussions to patients when providers lack reliable current information about cancer care or fail to change practice patterns based on new evidence. Most alarming among these were misdiagnoses that caused lengthy treatment delays, resulted in unnecessary surgery or incorrect treatment, and jeopardized patient survival or quality of life. Speakers also noted that providers may miss signs of cancer in patients with other chronic illnesses or fail to detect depression or other serious health problems in cancer patients. Provider education in these areas, on the care and needs of dying patients, and on cultural differences that affect care were listed among the areas of greatest need in provider education.

Finally, the lack of effective medical data and reminder systems in most clinical settings was cited as a significant reason why some patients “fall through the cracks.”

As one physician observed, “Medicine currently is less computerized than Wal-Mart.”

Information and Education Barriers Faced by Patients and the Public

The lack of accurate cancer-related information that is readily available, understandable, clear, and delivered in a sensitive and culturally acceptable manner is a major contributor to the inability of patients and the public to obtain the most appropriate cancer prevention, treatment, and supportive care.

Cancer is perhaps the most feared of diseases. As speakers described vividly, fear of cancer is shared by virtually all
populations and cultures, and takes the form of fatalism, fear of treatment and its costs, fear of pain and disfigurement, and fear of rejection by one’s partner, family, or community. These fears, and enduring myths about cancer, cause many people to reject cancer prevention messages, avoid cancer screening, delay or avoid treatment if diagnosed, and have difficulty understanding and choosing among treatment options or following treatment regimens.

For some Americans, cancer-related information is simply unavailable. Thousands in remote regions and high poverty areas lack telephones and in many areas library access is limited. Some remote areas even lack radio or television reception. Few of the poor own computers or have Internet access. In addition, many people do not go to the doctor or other sites where cancer information is most commonly found. Reaching these populations is a major challenge that some communities are addressing through outreach efforts, church-linked activities, and collaborations with local agencies that provide financial or other assistance.

For many other Americans, available cancer information is unusable due to literacy, language, or cultural barriers. Low literacy appears to be an underappreciated barrier to cancer information and care. Language barriers are a growing challenge to effective cancer communication as the current wave of immigration continues and are a special problem in the clinical setting, where relatives, including children, or strangers are being called upon to translate detailed information about test results or treatment. Speakers reported that some written materials mistakenly have been translated at too high a reading level or in the wrong dialect. Moreover, in some languages, no translation is possible; for example, in some Alaska Native languages, there is no word for cancer. Cancer information also must be presented in a culturally sensitive manner if the messages are to be accepted.

Cancer survivors and health care professionals emphasized repeatedly that regardless of educational level, income, or insurance status, people need help finding and evaluating accurate, up-to-date cancer information and navigating the complex and fragmented health care system. Communities are attempting to address this need by training community members to be outreach workers and cancer educators and by establishing “patient navigator” programs in hospitals and other treatment facilities to help people access medical and financial assistance for which they may qualify and secure the care they need. While the need for such programs
for insured and uninsured patients at all educational and socioeconomic levels was unquestioned, the tenuous stability of these often fledgling programs was also underscored.

Finally, speakers indicated that cancer surveillance is grossly underfunded in many areas of the country. Without adequate information on the extent and nature of the cancer burden, states cannot identify high risk groups, focus their planning efforts, develop targeted prevention and cancer control efforts, or evaluate their success. Local data on cancer patterns and trends may take years to accumulate, but these activities should proceed in tandem with action to address readily apparent cancer control problems.

The Impact of Culture and Bias on Cancer Care

Disparities in cancer treatment and disease outcome between various population groups are being documented with increasing frequency and clarity. A considerable number of speakers recounted experiences in which they or others received—or did not receive—cancer information or care for reasons stemming from cultural or racial differences, and biases these differences engendered. Importantly, bias that results in negative health outcomes can originate from both patients and health care providers.

Issues of Culture and Bias Originating With Patients and the Public

Cultural perspectives or biases may cause individuals to avoid cancer screening or treatment, or otherwise make decisions that may adversely affect their survival and quality of life. These biases can also have a positive impact on health. They affect the ways in which people perceive illness, how they develop and act on medical and caregiver preferences including folk healing methods, how they explain and tolerate pain, and what they perceive to be quality care. As numerous speakers indicated, however, fatalism about cancer remains pervasive in many cultures, though it takes different forms in different cultural groups. Old myths about cancer also persist in many populations.

The grinding circumstances and resulting culture of poverty profoundly affect the information and care-seeking behaviors of the poor. Rural residence and agricultural lifestyle also comprise a distinct culture in which it is rare to seek medical care unless one is in significant pain. Many rural residents, some of whom are poor, are uninsured and do not believe in
going to the doctor unless they can pay the bill. In addition, farming women are unlikely to interrupt farm routines to seek medical care, even when they have symptoms. Similarly, speakers suggested that the cultural importance of fulfilling one’s role in the family causes many Asian women to minimize their own health needs and avoid out-of-pocket health care expenditures.

Provider-patient relationships built on familiarity and trust are crucial to effective education and medical care for some populations, notably Native Americans and a number of recent immigrant populations. Yet the medical facilities at which these populations typically receive care are staffed by temporary duty doctors who often are of different cultures or lack sensitivity to the cultures of their patients. In some cases, resistance to entering the health care system and difficulty navigating it are undergirded by cultural traditions that consider assertiveness, particularly with authority figures, to be inappropriate or rude. Speakers testified to the critical need for health care providers from minority and underserved populations. In addition, traditions concerning female modesty and the acceptability of female patients being examined by male physicians underscored the need for more female health care providers.

Distrust of the health care system generally is common, particularly among populations that historically have been targets of discrimination. Many people fear being used as “guinea pigs” by medical practitioners. This distrust reinforces the fear of cancer treatment, including clinical trials, and remains a significant barrier to appropriate cancer care.

Secrecy about cancer remains prevalent in some populations, with patients hiding their disease even from partners and other family members. In some Asian cultures, knowledge of a cancer diagnosis still is withheld from the patient. These cultural prohibitions can make it extremely difficult to reach people with needed cancer information and care.
Issues of Culture and Bias Originating With Health Care Providers

Bias, either overt or unintended, also can originate with health care providers and administrators who may make decisions or recommendations that are not in the patient’s best interest. Physicians in particular have a special responsibility to be sensitive to their own cultural mindsets and biases, as well as those of their patients, because of the power and authority many patients confer upon their doctors. Speakers testified to pervasive and often overt provider bias against gay and lesbian patients that causes these patients to avoid screening and care. Patients with disabilities may experience unintended bias when they are not offered screening or other cancer-related care because providers focus only on health issues related to the disability. In other cases, providers may not share clinical information with patients who they assume will not understand it, or may fail to offer treatment regimens they assume patients will be unable to follow.

Numerous speakers indicated that bias at the provider and institutional levels also may occur when assistance, referrals, treatment, and other services are more readily offered to more educated or white patients compared with less educated or minority patients, even when they have equivalent resources, or lack of them. Some disparate behavior may be intentional, but according to speakers, more often reflects biases transparent to the providers themselves.

In addition, presenters described instances in which culturally insensitive behavior on the part of providers reflected a simple lack of education about other cultures and customs or an unwillingness to accommodate non-medical traditional practices that would not interfere with treatment but would comfort the patient.
What Can Be Done to Help People Now: Recommendations

The President’s Cancer Panel is acutely aware that the issues and problems described in this report are not being expressed for the first time. Indeed, the very fact that these problems—faced by real people with cancer every day, in every corner of the Nation—remain so prevalent makes concrete, achievable action to resolve them that much more urgent.

Access to appropriate cancer care is the crucial, fundamental step needed to relieve the desperate physical suffering, financial devastation, and loss of dignity so many people endure when cancer is diagnosed. If we lack the political will to craft and implement a National plan to address this unacceptable situation, then incremental steps must be taken to quickly remedy health care financing and delivery system elements that result in so much of the needless distress now experienced by cancer patients and survivors, and their families.

Continued research on the quality and equity of cancer care, outcome disparities, and related health economics and system issues is essential to guide transformation of the health care system in the coming years to better serve the public. But the people with cancer today, and their families, cannot wait for this distant relief. The President’s Cancer Panel recommends:

Immediate Action Steps

1. Provide immediate medical coverage for the uninsured—84 percent of whom are workers and their dependents—upon a diagnosis of cancer to help ensure that no person with this disease goes untreated.

2. Address health coverage issues that contribute substantially to the financial devastation of people underinsured for cancer care costs:
   - Provide reimbursement for anti-cancer agents, supportive medications (e.g., antiemetics, pain medications), and proven chemopreventive agents, regardless of method of administration.
• Within two years, public and private payers should reach consensus on and implement a standard health benefit package for cancer care. This benefit package should be based on the best available medical evidence and should be updated regularly to reflect advances in the standard of care. The reports and deliberations of the Institute of Medicine, other groups, and consumers should be used to inform this effort.

3. Address patient and public needs for cancer information and for assistance in accessing services:

• Provide funding to help communities coordinate, promote, and support community-based programs, including patient navigator programs, that help people obtain cancer information, screening, treatment, and supportive services.

• Recognize that the services of non-physician personnel who are trained to conduct cancer screening, and provide cancer education and case management in varied settings are an important component of cancer care that should be reimbursed.

4. Sustain cancer care in the community by providing consistent and realistic health care provider reimbursement across states, and between urban and rural locations within states, for the cost of chemotherapy drugs and their administration.

Longer-Term Solutions

1. Medicare, Medicaid, the Veterans Administration, the Department of Defense, the Indian Health Service, and other public payers should clarify the order of responsibility for payment for cancer care services when individual patients are eligible for benefits under more than one program. This information should be communicated promptly and clearly to those who provide cancer care services and assist patients in navigating the health care system. The existing Quality Interagency Coordination Task Force may provide a forum for accomplishing this important task.
2. Develop Federal policies to minimize bias in the provision of cancer care:
   - Raise awareness of unintended or overt bias through initial and continuing training of health care professionals at all levels, as well as administrators and others who make decisions affecting patient care.
   - Establish and implement systems for monitoring treatment equity. In addition, expand quality of care research to include issues of treatment equity.

3. Minimize disparities in the provision of cancer care by:
   - Educating primary care providers about cancer.
   - Educating all cancer care professionals about the nature and application of evidence-based medicine and about clinical trials.
   - Developing and disseminating better tools to assist health care providers in conveying information about cancer and about cancer care options.

4. Address the problems of temporary medical staffing and cultural incompatibility by establishing additional mechanisms to encourage more minorities and members of other underserved populations to enter cancer care professions. Provide incentives to encourage providers to practice in medically underserved areas.

5. Extend state-of-the-art cancer care to rural, frontier, and other underserved areas by expanding the use of telemedicine and providing a reimbursement system that facilitates expansion of telemedicine to geographically underserved areas.

6. Permit more flexible use of categorical funding where appropriate to enable states to fashion more rational and more comprehensive cancer control programs.
The President’s Cancer Panel is charged under the National Cancer Act of 1971 to monitor the development and implementation of the National Cancer Program and to report promptly to the President on barriers to the Program’s effective implementation.

In 1999, following its report on issues of cancer care quality and cancer-related quality of life,1 the Panel evaluated the evolution and current status of the National Cancer Program as a whole, including its research and delivery components—both public and private—as well as some of the social, economic, and information-related factors that influence its effectiveness. That report to the President described a serious dissonance between the cancer care that research evidence has proven to be effective and the extent to which this care is provided to Americans with cancer and those at risk for the disease.2

These findings were the impetus for a series of seven regional meetings conducted by the Panel during 2000 and 2001 to explore in greater detail the barriers that are keeping all Americans from receiving the most appropriate cancer care. Each regional meeting brought together representatives from each of seven to nine states, the District of Columbia, and U.S. Territories (see Appendix A). At the conclusion of the seven meetings, testimony had been received from every state and territory in the nation. Those offering formal testimony included cancer survivors, family caregivers, State health department representatives, local and regional public and private program administrators, health care providers at all levels, advocates, and volunteers. In addition, all but one meeting included an evening Town Hall to provide additional opportunity for local citizens to raise issues and share personal experiences with the Panel.

In all, 393 individuals provided testimony; of these, 163 were cancer survivors. The Panel appreciates the commitment of so many of the speakers who traveled across great distances, left jobs and family, and endured their own hardship or personal pain to describe the issues of cancer care in America at their most individual levels. The Panel is profoundly grateful to the following individuals who devoted a part of the limited time remaining to them to tell their stories, and who have since died:

- Dante Delledonne
- Ken Giddes
- Karen Kitzmiller
- Sue Kocsis

The following report and recommendations are based on all of the testimony received from June 2000 through May 2001. The Panel recognizes that invited testimony does not carry the weight of empirical study. However, the Panel believes there is a point at which anecdotes become a body of qualitative evidence, and the importance of the issues echoed repeatedly and consistently throughout every part of this country cannot, and must not, be dismissed. These voices of a broken health care system must be heard.
Who Has Cancer in America?

The mother of your child’s best friend. The man who waited on you at the hardware store last week. Your brother-in-law. The woman in front of you in the check-out line at the market. Your favorite teacher in high school. The homeless woman who’s always at your subway stop. Your doctor. The eight year old son of the family three doors down. Your co-worker at the plant who brings doughnuts for everybody. Your minister. The teller at the bank. Your college roommate’s wife. The farmer who grew the potato you’ll have for dinner tonight. Your grandmother at the nursing home. The father of your paperboy. Perhaps, even you.

Right now, 8.9 million people in America are living with cancer. They will be joined by more than 1.2 million others who will be newly diagnosed this year—about 34,000 each day. Overall, cancer is an equal opportunity disease—it strikes people of all ages, races, ethnicities, religions, cultures, geographic locations, and education and income levels. It strikes both the influential and the invisible among us. It kills more than a half million of us every year. And every person is at risk for cancer in his or her lifetime.

At the same time, certain cancers are known to be more or less common among specific populations, for reasons that may include varying combinations of lifestyle behaviors, genetic predisposition, and environmental exposures. For example, lung, bladder, esophagus, and oral cancers are most common among current and former tobacco users. Skin cancers are more common among fair skinned people who have had excessive sun exposure. The chance of developing most adult tumors rises with increasing age, seemingly due to accumulating genetic errors, the aggregate effect of environmental exposures, and other factors. Neuroblastoma and Wilm’s tumor are among the cancers that occur almost exclusively in children and are believed to be due to inherited or very early gene mutations. It is less clear why the prostate cancer rate among African American men is 60 percent higher than that of white men, why Vietnamese women have exceedingly high cervical cancer rates, or why nasopharynx and stomach cancer rates are unusually high among some Asian populations. Nor do we know why some Latino, American Indian, and other populations have lower than average rates for certain cancers.

Modest decreases in cancer incidence and mortality have been achieved in recent years, yet cancer remains the second leading cause of death in the nation, and is responsible for immeasurable fear, suffering, and hardship among both patients and their families.
quality of life. But many of the interventions that research evidence has shown to be effective are not reaching people throughout the country. The Panel illustrated this “disconnect” between the cancer research and cancer care delivery efforts (see Appendix B) in its 1999 report on the evolution and status of the National Cancer Program, and emphasized that this disconnect is not a failure of the research effort. Without question, the cancer research effort has been productive, and continued research is needed, both to expand basic research and to translate basic science discoveries into useful interventions; to improve cancer preventive, palliative, and end-of-life care; and to enhance our understanding and measurement of cancer care quality, including cancer communications effectiveness, treatment outcomes, and quality of life. But the current disconnect reflects the nation’s ongoing failure to organize, finance, and operate a health care system that assures access to appropriate cancer care and brings proven interventions to all of the population. In short, our health care system is broken, and it is failing people with cancer and those at risk for cancer—all of us.

The Health Care System and Cancer—Who Is Underserved?

The National Cancer Institute’s (NCI) Special Populations Working Group has defined the term “underserved” as referring to “populations at risk of poor physical, psychological, and/or social health who experience a lack of sufficient community, clinical, or individual resources to effectively meet their needs.” This definition encompasses groups that are not benefitting from available options for improving health and reducing disparities in the United States. In this sense, underservice is a dynamic state into which an individual may move in or out, and varies by type of disease and disease stage. The term “medically underserved,” refers more narrowly to Federally designated geographic areas with poor access to health care providers and medical services.

Thus, while the underserved most often are thought to be the poor, minorities, the uninsured, and those in medically underserved areas—populations that indeed are represented disproportionately among the underserved—a person also may be underserved for cancer care even though he or she has insurance, if that coverage does not provide needed services at a time of health care crisis, or to prevent and detect disease. Moreover, even those with comprehensive insurance may be underserved if their health care provider lacks information about the most appropriate evidence-based cancer care, including clinical trials.

Health Care System Changes

Although the fee-for-service (FFS) system that dominated health care financing prior to the advent of managed care was a major contributor to spiraling health care costs, changes in the structure and financing of the health care system in recent years have had profound effects on the quantity and scope of cancer care provided, provider payments for care, and the system’s capacity to care for the indigent. Much of the change has been due to cost containment pressures from managed care plans and “managed” FFS plans that are the payers for a growing percentage of the population in both private and publicly funded insurance settings. “Gatekeepers” in many health plans restrict referrals to specialists, exclude the use of drugs not on a health plan’s formulary, limit supportive and rehabilitative care, deny access to clinical research trials, refuse to pay for the routine care costs of those who participate in trials, and set strict limits on payments for medical equipment and prostheses.

In addition, reimbursements to providers of all types have fallen steadily. To compensate, physicians in office and institutional settings are under pressure to see more patients in less time. Physicians in office settings may suffer financially or be dropped from a health plan’s provider roster (thereby losing patients) if,
in the judgment of the plan, they spend too much time with individual patients, make too many referrals, or order too many tests. State-of-the-art cancer care is increasingly expensive and technology driven, but with reduced profits and reimbursements, physicians and institutions, particularly those other than major metropolitan medical centers, are finding it increasingly difficult to buy the equipment and supplies needed to provide the most effective tests and therapies. This legitimate need to upgrade cancer care technology for the benefit of patients is to be distinguished from unnecessary duplication of resources often driven by competition between providers, particularly in urban areas. Reimbursement reductions also are affecting hospital staffing patterns (particularly for non-physician personnel), the willingness and ability of institutions or individual physicians to provide care for those unable to pay, and the ability of cancer patients to receive second opinions and appropriate care alternatives.

For many patients, these health system changes have resulted in a fragmented, often bewildering system in which cancer and other health care must be obtained only at facilities and from providers under contract to the health plan. In this setting, continuity of care may be compromised, patients may no longer have regular and trusted providers, supportive services are minimal or non-existent, and responsibilities for patient education and appropriate sharing of medical information frequently fall to patients themselves. For some, there simply is no care available.

**Distribution of Cancer Care Resources**

People living in rural, frontier, geographically isolated, and impoverished inner city areas suffer the most from the uneven distribution of cancer care resources and providers—a key element of underservice in these populations. This uneven distribution of resources and personnel is longstanding, and may be worsening as reimbursement changes make it less and less attractive for providers to establish and maintain practices in rural and inner city areas. Provider reimbursements for most aspects of care are lower in rural areas compared to metropolitan settings. Facility and certain other overhead costs are lower in rural areas, but the reimbursement differences appear to exceed substantially these differential costs of providing care. With sparse populations in rural and frontier areas, providers cannot sustain operating revenues or accumulate capital to buy new equipment and supplies by seeing more patients.

**The Underinsured**

In the current health care environment, simply having insurance is no assurance of appropriate care, particularly for complex diseases such as cancer for which treatment is expensive and often lengthy. At least 31 million insured Americans are estimated to be inadequately insured for a catastrophic disease such as cancer.

This estimate of the underinsured does not include Medicare beneficiaries, a large percentage of whom live on fixed incomes. Many cannot afford Medicare Part B premiums (for outpatient and all physician care) or Medicare supplemental insurance policies, and so are at risk for all health care expenses incurred outside of a hospital setting. At this time, outpatient prescription medications are not covered under Medicare; these can total several hundred dollars per month even for patients who do not have cancer. Many elderly Americans routinely must choose between buying food or their medications. Some report reducing their dosages, only buying part of a prescribed amount of medication, or sharing prescriptions with family members or friends.

Lower-income employed people who can afford to participate in employer-sponsored group insurance plans often choose limited coverage with high deductibles and copayments in order...
to lower premiums to affordable levels, but then find their coverage inadequate (both in terms of non-covered costs and access to the most appropriate care) when diagnosed with a serious disease such as cancer. These families find they must deplete their savings and sell their homes and other assets to pay the non-covered costs of cancer. Middle-income people can likewise be devastated financially by the costs of cancer care, though they initially may have more resources, including family support, on which to draw. Additionally, many people with cancer lose their health coverage when either the patient or spouse loses the job through which insurance was offered. Many cannot afford to continue coverage under provisions of the Consolidated Omnibus Budget Reconciliation Act (COBRA)\textsuperscript{11} that enable such workers to maintain their health insurance under an employer’s group plan if they pay the full monthly premium.

People with Medicaid or other publicly funded medical assistance typically must accept whatever care is offered since they have no income with which to purchase private insurance or pay out-of-pocket for additional care. Some populations are more likely to rely on public insurance; more than one-fifth of Hispanics and over one-fourth of African Americans have public insurance only (including Medicare), compared to one-tenth of whites.\textsuperscript{12} In addition, many cancer care providers hesitate to accept Medicaid patients because of low reimbursement rates and slow payment processing.

The Uninsured

Currently, 44 million people in America have no health insurance at all,\textsuperscript{13} and therefore little access to health care services, including cancer screening and cancer care. When they receive care at all, this population tends to rely on special screening programs, charity care, and care provided by hospital emergency departments. According to a 1999 report of the National Coalition on Health Care, under favorable economic conditions the number of uninsured non-elderly (those under age 65) is projected to rise to 52 to 54 million by 2009; if a recession occurs, that number will likely jump to 61 million.\textsuperscript{14} An analysis of the 1999 Current Population Survey (CPS) indicated that in 1998, ten states had uninsured rates in excess of 20 percent of their non-elderly population, and the uninsured rate among the non-elderly nationwide was 18.3 percent.\textsuperscript{15} Uninsured rates also appear to be affected by the availability and eligibility requirements of state-level Medicaid or other medical assistance programs. Data from the National Center for Health Statistics (NCHS) indicate that an important factor in uninsurance trends is the decline in private insurance coverage, particularly for people aged 18 to 44 years, whose rate of coverage fell from 77 percent in 1984 to 69 percent in 1997.\textsuperscript{16} A recent study indicated that near-elderly (aged 55 to 64) working women with health problems are more likely than younger workers to lack employment-based health insurance, to be uninsured, and to lack sufficient income to purchase insurance.\textsuperscript{17} Changing employment trends appear to be exacerbating the problem of uninsurance. Many larger employers are cutting the size of their full-time work force and replacing these employees with part-time or contract workers who typically do not receive health benefits.\textsuperscript{18} More people are now working at small companies that cannot afford to offer health coverage. Fewer employers of all sizes are offering health insurance. A growing number of the uninsured appear to be former welfare recipients who have moved into the workforce. Though no longer eligible for Medicaid, they do not have health coverage through their new jobs.\textsuperscript{19} Nationwide, 84 percent of the non-elderly uninsured are working adults and their children.\textsuperscript{20}

The CPS data analysis also indicates that while 53 percent of the uninsured are non-Latino whites, ethnic minorities have disproportionately high uninsurance rates—38 percent of Latinos, 24 percent of African Americans, and
22 percent of Asian American/Pacific Islanders are uninsured, compared with 14 percent of non-Latino whites.21

Evidence of Unequal Treatment

When a person has a diagnosed or suspected cancer—any person, with any cancer—he or she needs immediate help. That means complete and understandable information, and the most appropriate care. But some people with cancer, even the same type of cancer at the same stage of disease, fare better than others for reasons that have little or nothing to do with the biological aspects of their disease. Growing evidence indicates that much of the disparity in cancer outcome has to do with the type, timeliness, and continuity of cancer care that people do or do not receive.

The impact of poor quality and unequal treatment on specific patient groups is being documented with increasing frequency and clarity. For example, a recent review of more than 50 studies of colorectal cancer treatment and outcome found that older, minority, and poorer patients tend to receive substandard care and have less favorable outcomes than those who are younger, white, or more affluent.22 Other studies have found that African Americans are less likely to receive potentially curative surgery for early stage lung cancer23 and less than the minimum expected care for breast cancer24 compared with white patients having equivalent income, access, and stage of disease. Both men and women of higher income are more likely than those with lower incomes to obtain cancer screening25,26 that can result in earlier detection of disease and in many cases, improved survival. Similar disparities among population groups have been documented with regard to non-cancer illness, including prescriptions for and access to pain medication,27,28 use of cardiac catheterization following a heart attack,29 and managed care plan approvals for emergency room care.30

Efforts to address disparity and quality issues also are accelerating as awareness and documentation of these problems increases. The Department of Health and Human Services (DHHS) is pursuing an initiative to eliminate health disparities by 2010, and individual DHHS agencies have been charged to implement efforts in support of this goal. For example, the DHHS Office of Minority Health has published final recommendations for culturally and linguistically appropriate service (CLAS) standards for health care.31

Specific to cancer, NCI has established a Center to Reduce Cancer Health Disparities32 and also has established research programs to address quality of cancer care issues and to advance outcomes research. The President’s Cancer Panel,33 the National Cancer Policy Board,34,35 the Institute of Medicine,36,37,38 the National Cancer Advisory Board,39 and others40,41,42,43 likewise have studied and continue to explore issues of cancer and other health care quality, equity, and outcome. All of these efforts, together with the accumulating body of qualitative evidence, a part of which is contained in the next section of this document,44 can be used to educate and guide policymakers to create a health care system that better and more equitably serves the public.
What Is Happening to Real People: Findings

At each of its seven regional meetings, the Panel posed the question: “Why don’t all Americans get the best available cancer care?” The numerous issues, barriers, and contributing factors raised in testimony provided to the Panel by cancer patients and survivors, family caregivers, representatives of state health departments, state and local program administrators and staff, physicians, nurses, hospice providers, outreach workers, advocates, and volunteers fall broadly into two categories—access and information. In addition, behaviors and decision-making by both health care providers and the public that stem from perceived differences among populations and individuals can influence greatly the type and quality of cancer care that people receive.

What Barriers Limit or Prevent Access to Care?

Access barriers include those related to the organization and operation of the health care system itself, financial barriers to care, and physical barriers that reduce or prevent access. These categories are not always mutually exclusive, however. For example, certain barriers that appear to be related to physical distance from care may be mediated by financial factors that have influenced the distribution of health care facilities, resources, and personnel. Speakers at the regional meetings and town halls across the country attested to the complexity and interrelated nature of these cancer care barriers.

System and Financial Barriers

System and financial issues are inextricably linked. Fiscal concerns have shaped, and continue to shape, both the private and public components of the health care delivery system.

“The health care system and reimbursement system is terribly broken...it’s incentivizing all the wrong behaviors. Preventive care is not reimbursable, really. Outreach is not reimbursable. Procedures are reimbursable...We need a system that fixes the basic overall funding of health care and that’s the key.”

Sharon Anderson, administrator, Delaware

“It is time for the nation’s public health work force to accept responsibility for leadership and coordination in the war on cancer. They should be the hub of activity at state and local levels. Voluntary groups cannot do the job alone.”

Lois J. Hall, screening project director, Ohio
Issues Related to the Structure and Focus of the Health Care System

Speakers from across the country, including program administrators, providers, and patients, emphasized that the current health care system focuses on acute care over prevention and allocates cancer funding by disease site. Both approaches, they stated, are counterproductive to the provision of comprehensive cancer care and the development of effective cancer control programs.

Currently, public health resources and leadership are insufficient to create consistent community goals and programs for cancer control, including cancer education and cancer prevention. Perhaps the most glaring example in this regard concerns tobacco control; while some states have devoted significant resources to this effort, other state programs are minimal. Early detection programs for cancer are likewise uneven. Nearly every state has a CDC-funded Breast and Cervical Cancer Detection Program, and in some states, adjunctive local programs targeting these cancers. Testimony heard across the country made clear that without Federal funding, few states attempt to screen for other cancers.

It also was noted that although the breast and cervical cancer screening programs exist, most are reaching only a small fraction of eligible women. A variety of factors, including funding restrictions and cultural barriers, appear to be responsible for such limited success. States have struggled with the problem of finding funds and providers to pay for the treatment of people whose cancer is detected through Federal government-sponsored screening programs. The Breast and Cervical Cancer Treatment Act of 2000 now provides supplemental Medicaid funding to pay for treating women whose breast or cervical cancer is detected through the CDC-funded screening effort, however, states must apply for and match these funds and some states may elect not to do so. Moreover, people whose cancer is detected through other screening initiatives may still find themselves with diagnosed cancer and no source of treatment.

Several state health department representatives described the difficulty of trying to fashion comprehensive cancer control programs with piecemeal and often tenuous funding. If a state or Federal grant for a particular service program is not renewed, those services are likely to be discontinued unless alternative funding can be found. In addition, each grant comes with its own documentation requirements and restrictions on how the funds may be used, even when multiple grants are from the same agency, such as the CDC.

“Every person who dies while we’re looking for care for them, or for whom we find care too late to make a difference, only makes it more difficult for us to convince the family and friends of the benefits of screening.”

Mawuna Gardesey, administrator, Delaware

“Assistance programs seem to be based on the political popularity of the specific disease...”

Susan Garrett, clinical coordinator, Maryland

Once the diagnosis of lymphoma was established for Lucinda, I spent weeks calling programs that offer assistance to people with cancer. One program only assisted people with leukemia. Another, only people with breast cancer. How does one say to a patient, ‘If you had this type of cancer, I could get help for you but I can’t get help for your type of cancer?’

Susan Garrett, clinical coordinator, Maryland
These restrictions often prevent states and localities from organizing and delivering services in the most logical and cost-effective manner. A few states have funding designated for a comprehensive cancer control program, but these relatively new efforts are not yet well established. Other states indicated that while they have or are developing cancer control plans, no funding has been committed to implementing these plans.

Intervention, demonstration, and other research projects funded by the Federal government, academic centers, foundations, or other private sources further complicate the picture by providing needed services for the three to five year period of the grant, only to disappear when the grant period concludes because provision is seldom made to sustain the program in the community once it has been shown to be beneficial. The community is abruptly faced with a gap in much-needed care, often just as residents have come to trust and depend on the providers of these services. This grant-based funding also hampers efforts to recruit and retain cancer care providers, particularly in underserved areas.

Lack of coordination among public payers was another issue cited repeatedly in discussions of how the health care system itself creates barriers to care. Some patients are eligible for cancer care coverage through more than one program, such as Medicaid, Medicare, the Veterans Administration (VA), and the Indian Health Service (IHS). Administrators of these programs do not appear, as a matter of course, to assist patients in determining how best to access all of the coverage to which they are entitled. In too many cases, according to testimony provided to the Panel, patients are treated like “hot potatoes,” with payers seemingly competing to avoid paying for care. The most consistent examples described involved the IHS; according to a member of the Cheyenne tribe in South Dakota, “they make sure that when we’re ill and we come to them, that they are a payer of last resort and that we understand that.” Similarly, a physician related the case of an older veteran who was encouraged to use his Medicare benefits instead of his veteran’s benefits to cover his cancer care. Unable to afford the Medicare Part B premium for all physician and outpatient care, he unwittingly paid out-of-pocket for services that would have been covered under the VA system. Even with Part B coverage, he still would have been subject to a 20 percent copayment not required for the same services under the VA benefit program.

“I had two female friends who in their late 40s died of cancer...both of their husbands were physicians...we complain that poor people sometimes don’t get adequate care, and yet these people were neither poor nor uneducated and they really didn’t get the care they needed. Even the insured and the privileged are not immune to the delay or the misdiagnoses and death.”

Mary Ann Andreis, family caregiver, Rhode Island

“I refinanced my home, which was paid for...I had to sell one of my cars. And it just bankrupted us. I didn’t have any more money. The copayments were so high it ate me up...Yes, I had Medicare and I have Medicare supplement. I’m very grateful that we have. I don’t believe in charging anything that I can’t pay. After I used up all my money then I [felt] like I have to have some help. So I went to Medicaid...I was making too much money. So you have to be almost starving to death before you can get Medicaid.”

Alan Hebert, prostate and colon cancer survivor, Louisiana
For people with employer-sponsored or other private health insurance, fragmentation of care, gatekeepers who control access to screening and specialists, and limitations or exclusions on specific drugs and services, including clinical trials, were the system-related barriers to cancer care most often described. Insured patients detailed situations in which they were denied care despite suspicious symptoms and/or a strong family history of cancer because they did not fit screening guidelines or the typical patient profile; this was particularly true for younger patients. Those with diagnosed cancer recounted having to battle their health plan continually to gain access to specialists, medications, and testing they needed. In some cases, physicians did not know the most appropriate care for a patient’s disease.

Several speakers emphasized that the current system discourages appropriate end of life care. Some public payers (e.g., IHS, VA) have hospice benefits, but according to speakers, do not refer patients to hospice or have insufficient service capacity for the beneficiary population. Medicare beneficiaries can receive hospice benefits only if their life expectancy is projected to be six months or less. Physicians testified that it is exceedingly difficult to predict how long some patients with advanced disease may live.

Many physicians wait too long to refer to hospice, fearing that if they refer too soon, the patient may outlive the benefit period. As a result, many dying patients and their families are deprived of much of the comfort and care that hospice could provide. Another system barrier to hospice care is Medicare’s requirement that the patient must have a primary caregiver living in the home; those who cannot meet this requirement often are forced to die in institutional settings. In addition, patients can receive regular Medicare benefits or hospice, but not both. Under Medicare, hospice patients may not receive palliative chemotherapy or radiation; hospice providers emphasized that many patients could benefit greatly from care that controls pain and other symptoms but does not have a curative intent or prolong life.

Financial Issues Affecting Patient Access to Care

Throughout the testimony in every region of the nation, from providers and patients alike, there was a consistent call for an equitable system of health insurance that would assure patients access to cancer preventive and therapeutic care when needed. Clearly, issues remain to be resolved as to what level of care should be considered the minimum...
expected care for specific conditions. However, numerous speakers voiced the same moral tenet:

No person in America with cancer should go untreated, experience insurance-related diagnosis or treatment delays that jeopardize survival, or be bankrupted by a cancer diagnosis.

For people with cancer and their families, health care system issues that can be explained logically by current fiscal and economic realities often translate into a personal reality that includes family bankruptcy, needless suffering, loss of dignity, and loss of life. Inadequate insurance, or a total lack of health insurance, were the financial barriers to cancer care most frequently experienced by testifying survivors and most often noted by state health department and local program officials. State health officials reported that uninsured rates are as high as 25 percent in some states for the non-Medicare population, with some states reporting markedly higher uninsured rates in rural and frontier regions compared with urban areas. The underinsured population is difficult to quantify in most states.

Many people discover after a cancer diagnosis that their employer-sponsored or individual private health insurance does not cover many of the costs of cancer care. Those with only catastrophic coverage typically have extremely high deductibles, but even those with fairly comprehensive coverage or Medicare find themselves faced with significant, unanticipated out-of-pocket costs. As numerous survivors and family members recounted, Medicare and many private health plans do not provide reimbursement for orally administered chemotherapy drugs; supportive medications such as those to control pain, nausea, and anemia; or chemopreventive drugs. Such out-of-pocket costs, plus deductibles, copayments, and the costs of laboratory and other tests, office visits, supportive care, and other denied or excluded services add up quickly.

Some providers demand full payment at the time of service, and slow reimbursements from the health plan strain cash flow even further. For example, a mother whose son died of leukemia recounted being asked by her local pharmacy to charge thousands of dollars a month on her credit card to pay for his medications because the pharmacy was unwilling to wait for reimbursement by the insurer. Such costs constitute an especially heavy burden for patients whose income is reduced or lost; even more costs are incurred by a spouse, significant other, or parent who is caring for the patient.
and consequently must take time off from work. These and other non-covered costs, such as transportation, child care, and medical supplies can rapidly drain personal and family savings, causing some patients to sell whatever assets they may have, including the family home, farm, or business, or to go without care.

Should a patient or spouse lose the job providing health insurance, the family can face a quick descent into indigent care and family bankruptcy. To qualify for Medicaid, patients may be forced to give up any remaining assets, including the car needed to reach the hospital for treatment. If they are well enough to return to work after treatment but have had a break in insurance coverage in excess of two months, cancer survivors may find they are subject to a waiting period of up to one year for coverage of pre-existing conditions under a new employer’s health plan. Survivors or family members (e.g., a spouse with cancer) who return to work at jobs that do not offer health insurance may be able to obtain private coverage, but often only at prohibitively high cost, and in some cases, excluding coverage for any services related to cancer. For example, a breast cancer survivor testified that she lost her insurance when she and her husband divorced; the only insurance she has since been able to secure excludes coverage for care in any way related to breast cancer, including screening mammography.

The unemployed poor, the working poor, and self-employed middle-income people are in an even more precarious position, since they are unlikely to have any health insurance at all. The poorest poor are the most likely to qualify for Medicaid or obtain care through hospital charity care systems. However, speakers described such long delays in obtaining Medicaid approval that they were faced with more advanced disease by the time they were able to begin treatment. In one such case, a cervical cancer survivor recounted beginning her quest for Medicaid approval when her disease was at a precancerous stage; by the time she was approved, she had advanced invasive cancer. Moreover, she lost her Medicaid eligibility as soon as her initial treatment was completed, leaving her with no coverage for needed follow-up care. Other survivors indicated that although they had secured a Medicaid card, it took weeks or months to find a provider who would accept Medicaid payment and provide needed care.

Many of the cancer survivors and family caregivers providing testimony were self-employed—farmers, ranchers, small
business owners, and other independent workers. As these speakers indicated, they seldom can afford even basic health insurance, though they make enough money to support themselves and their families. The working poor may hold two or three jobs, none offering health benefits. Both groups are likely to lack both health insurance and the financial reserves to see them through an extended illness. Upon being diagnosed with cancer, however, these patients often find that they make too much money to qualify for Medicaid. If they stop working in order to become eligible for Medicaid, they may not have enough income to support themselves and their families. This wrenching predicament was described by many of the survivors providing testimony to the Panel. One such case involved a self-employed cancer survivor from the District of Columbia, who was the sole source of support of her elderly mother. Though her mother was a Medicare beneficiary, her medications, costing $450 to $3,500 per month, were not covered by any insurance. Despite a family history of cancer, the daughter had chosen to forgo health insurance for herself in order to pay for her mother’s medicine. When she was diagnosed with cancer, the daughter applied for Medicaid but was denied because her income exceeded the maximum allowable amount. Yet she could not give up her income to qualify for Medicaid without depriving her mother of needed medication.

People with cancer who are unable to work due to their disease may qualify for Social Security Income (SSI); however, one must have a diagnosed disease for at least 12 months before an application can be submitted, and it is quite common for applications to be denied two or more times before approval. Speakers reported that SSI income often exceeds state Medicaid income ceilings; the result is that patients may regain income to help meet living expenses, but lose their access to cancer care. Patients who are approved for Social Security Disability Income (SSDI) also become eligible for Medicare, but only after having a totally disabling disease for two years.

Knowing they cannot pay for care, some uninsured patients avoid screening that might result in a cancer diagnosis, and if diagnosed, refuse care in order to avoid what one presenter described as the “roller coaster” that leads to bankruptcy, or accumulating debts they fear might be left to family members to pay. It was suggested that those with incomes at 100 to 250 percent of the poverty level are most quickly wiped out financially by a cancer diagnosis.
Many uninsured patients struggle to pay out-of-pocket for their care, often amassing debt equal to several years’ wages, even when providers have discounted services and arranged for donated medications. To pay these bills, patients and family members take on second jobs, give up their homes and possessions, exhaust savings, and accept loans and donations from friends. A survivor of childhood cancer, now age 26, indicated that his family is still paying hospital bills incurred following his diagnosis at age ten. A man whose wife had cancer and died is working three jobs to pay more than $100,000 in bills for her care—though his wife did not survive, he felt she had received the best possible care and felt a responsibility to see that her doctor and hospital did not go uncompensated. Other patients whose creditors will not accept installment payments are forced to declare bankruptcy.

Financial Issues Affecting Health Care Providers

As health care payers and purchasers struggle to contain health care costs, financial pressures on health care providers continue to increase, affecting the care available to people with cancer and those in need of screening, diagnostic, and preventive services. Survivors and family members reported widely varied experiences with the provider community in terms of willingness to accommodate patients’ lack of insurance or full insurance coverage. Some patients were told “pay what you can, when you can;” others were able to negotiate reduced rates and payment plans with their hospitals and physicians; and some providers donated their time and services. In other cases, however, providers asked for advance payments of $20,000 to $100,000 before initiating treatment. These highly divergent responses to patients in need appear in some instances to reflect the extreme and intensifying conflict between some providers’ commitment to render care as needed and a range of financial pressures that threaten the solvency, vitality, ethics, and integrity of health care institutions and individual providers alike.

Providers at nearly every regional meeting reported steadily increasing payer-related paperwork, regulatory requirements, and other administrative responsibilities. Both institutional and office-based cancer care providers have been forced to hire staff whose sole job is to negotiate with payers to secure approvals for patient treatment and subsequently to try to obtain payment. These substantial administrative costs have been coupled with steadily falling reimbursement rates for drugs and services.

“Well, last month I come home from my chemo treatment and there was a message on my machine to call [the state of West Virginia] where I work...and the girl in the personnel office regretted to inform me that the state will no longer provide my medical coverage...I pay $600 a month now for my insurance to keep me going so I can receive my treatments. There’s no help available. I’ve called as many people as I can...I still have the Veterans [Administration] to go through yet because I’m a two-time Vietnam veteran...

We depleted our Christmas club and our savings account. I do receive Social Security disability. That’s $1,087 a month. When you subtract $600 a month from that, it doesn’t leave very much for food and bills and other things. And I also have one drug that I get 16 shots a month between chemo treatments, that’s for the white blood cells...and the copayment for that drug is $1,176...and my copayment for the pain medication is $130 a month. And my nausea medication, which I do have to take...five pills, my copay is $36. And sometimes those five pills run out in a week or sometimes they run out in less than a week.”

Edward Chisholm, colorectal cancer survivor, West Virginia
Many of the providers and advocates who testified before the Panel expressed grave concern about the effects that Medicare’s Ambulatory Payment Classifications (APCs, also known as the Hospital Outpatient Prospective Payment System, or HOPPS) will have throughout the cancer care system. For example, the cost of chemotherapy administration (e.g., nursing staff time, monitoring equipment, supplies) is not reimbursed in the office setting, and providers have previously covered these costs through margins on drug acquisition reimbursements. With severe drug cost reimbursement cuts, in some cases below acquisition cost, providers indicated they will have little choice but to absorb chemotherapy administration costs or stop providing office-based chemotherapy. As speakers pointed out, the solution to this problem is not necessarily reinstatement of higher drug reimbursements, but fair reimbursement for both chemotherapy administration and drug acquisition costs.

The hospital outpatient reimbursement scheme likewise threatens an already fragile balance. Many smaller community hospitals are going out of business, worsening access problems for rural patients. Even in large metropolitan areas, hospitals are struggling to maintain both services and solvency in the face of shrinking reimbursements by Medicare, Medicaid, and private payers. However, while demand for indigent care is increasing, decreased total revenues are reducing hospitals’ ability to absorb the direct and indirect costs of uncompensated care. Other casualties of these fiscal changes are nursing, pharmacy, social work, education, outreach, tumor registry, and other supportive care personnel, clinical research, and training programs for the next generation of cancer care providers.

Relative to differences in the cost of doing business, reimbursements for identical care are inappropriately inconsistent both across the nation and between urban and rural areas (even within the same state), with substantially lower reimbursements for the same items in rural areas. These payment differences have serious effects on the quality and type of care available in rural localities. As a radiation oncologist from the Midwest pointed out, “It costs the same amount of money for me to buy a linear accelerator as it does for somebody in New York City.” Radiotherapy and other complex medical equipment is extremely expensive and has a limited useful life as treatment techniques become more sophisticated. Supplies, such as the radiation sources required to operate specific pieces of equipment, may cost in excess of $30,000 and last only months—whether they are used or not.
Lower reimbursements for the same care provided in rural areas also has meant that instead of increasing efforts to take state-of-the-art cancer care to rural sites, many metropolitan cancer centers bring rural patients to the city for care. This approach constitutes a physical and financial hardship for many fatigued and ill patients, who incur travel, lodging, food, and child care costs to obtain care, especially if they must be accompanied by a spouse or companion. Oral medication, even if available for an individual’s condition, is not an alternative, since Medicare and many private plans do not cover the cost of “take home” drugs. Thus, some patients drive hundreds of miles just to receive an injection, then return home the same day.

Some providers are attempting to establish viable rural oncology clinic networks, but the combination of recently reduced reimbursements for drugs, lower patient volumes, and lack of reimbursement for chemotherapy supplies and administration are making financial survival questionable. The CEO/oncologist of such a network in Alabama described the situation this way: “We don’t have as many patients per nurse, per vial of Taxol®, per square foot of building, to dilute fixed cost...it doesn’t take long for the private sector to ratchet right down to many of the Medicare fee schedules, and the margins that are allowed are just unsolvable in the private doctor’s office.”

Another concern expressed was that Medicare APCs will reimburse for older drugs and treatment regimens, but not for newer anti-cancer agents and therapies. A related, ongoing issue is that Medicare-approved drug lists are not kept current, causing providers to engage in often lengthy negotiations with Medicare contractors for approval of newer treatments. Physicians reported instances in which paperwork and approval-related delays were so protracted that the patient’s disease advanced beyond the point at which the recommended treatment might have been curative, or even possible.

Speakers maintained that Medicare and Medicaid reimbursement cuts have “downstream” effects that are increasing the gap between state-of-the-art care and its implementation for all people. And as noted above, Medicare reimbursement levels and policies often are quickly adopted by the private health insurance industry. Reimbursement reductions under the APC payment schedule are of concern to all providers, including those whose patient population may be insured principally by private sector payers.
Testimony indicated that one effect of diminishing reimbursement levels is growing pressure on physicians in office and institutional settings alike to see more patients per day. Payers specify the maximum allowable reimbursement for office visits of various durations and the diagnosis codes that may warrant extensive office visits. Physicians who spend too much time with each patient risk a greater percentage of denied or reduced claims and may be dropped from health plan provider lists if they are judged to be inefficient. Physicians may also face penalties if they perform too many expensive tests or biopsies, even when these are deemed necessary.

While the effect on the provider may include reduced revenues and greater professional and personal stress, effects on the patient may include inadequate discussion of medical history and all relevant health issues; inadequate patient education; missed opportunities for preventive care; failure to be offered clinical trials or other care that requires lengthy explanation, monitoring, or additional paperwork; and in the worst cases, misdiagnoses.

Physical Barriers

Distance from Care

Testimony provided to the Panel suggested that residence in rural or frontier areas poses a special set of problems that comprise a vastly underappreciated barrier to cancer care access. It is estimated that one-fourth of the U.S. population of 281 million in 2000 live in areas designated as rural (i.e., fewer than 2,500 people per town boundary) and frontier (i.e., fewer than 6.6 people per square mile). In fact, roughly 45 percent of the U.S. land mass is frontier, and half of all states have frontier regions.

Excessive distance from cancer care is due in part to the concentration of health care personnel and resources in urban areas, the lack of public transportation systems in rural and frontier areas, and the fact that many residents of these areas lack the resources to travel to care.

Uneven Distribution of Cancer Care Resources and Personnel

The scarcity of health care providers and resources in rural and remote areas is a longstanding problem that Federal and State governments have tried to address over many
years (e.g., through the National Health Service Corps, rural and migrant health centers, and other mechanisms). Yet the number, type, and distribution of cancer care providers, treatment facilities, and treatment technology remains seriously inadequate in large areas of the country. In many underserved areas, there also are few primary care providers—in some counties, none at all. Where there are no primary care physicians, patients cannot get referrals to specialty care. As a result, people living in these areas frequently are misdiagnosed or diagnosed at late stages of disease. It was suggested that some cancers in these populations are never diagnosed.

Lower reimbursements for care provided in rural and frontier regions and the limited grant funding available in sparsely populated areas continue to hamper efforts to recruit and retain oncologists and other cancer care professionals and are a major contributor to the uneven distribution of cancer care services across the country. A young physician testified that he would like to practice in a rural setting, but feared he would not make enough money to repay his educational loans. Speakers called for programs and incentives to attract rural and minority individuals to cancer care professions and help them sustain careers in underserved areas.

A number of speakers noted that both oncology nurses and other nursing staff are in short supply nationwide and that their number is shrinking. The average oncology nurse in practice is 40 to 45 years old, and the number entering the field is decreasing because of health system changes. Most importantly, hospitals attempting to reduce costs in the face of shrinking reimbursements are replacing oncology nurses with nurses who are not specially trained to care for cancer patients. Apart from their specific training to manage cancer care regimens and treatment side effects, these nurses are critical to patient and family communication throughout the treatment process. Some nurses are leaving the profession because they are consistently being asked to care for too many patients at one time and these patients tend to be quite ill. In the current health care environment, inpatient care is reserved for the sickest patients and the most difficult treatment regimens. Concerns were expressed that such heavy caseloads may jeopardize patient safety. In addition, with fewer applicants, oncology and other nurse training programs are closing. Similar patterns are being observed among other cancer-related health professions. These trends bode ill for the future, as the elderly population and the
number of cancer patients and survivors grows, and cancer care becomes increasingly complex.

Some services cannot be taken easily to remote areas. One such service is radiation oncology. Providers testified that the lack of radiation facilities in rural and frontier areas results in more surgical treatment for breast and prostate cancer (due to both physician recommendation and patient choice) because the nearest radiation facility is too far away for patients to make daily round trips for several consecutive weeks. Physicians have tried to establish rural radiation facility networks, but these are proving to be barely self-sustaining. It likewise has proven difficult to sustain home hospice services in rural areas; a provider of these services indicated that a nurse may spend two hours traveling to make a one-hour home visit.

Speakers also noted that disabled patients face unique physical barriers to care. People with disabilities such as mobility, spasticity, and mental problems may require special equipment or specially trained personnel to enable them to receive cancer screening and care.

Transportation Issues

In rural and remote areas, there are virtually no public transportation systems. Patients reported driving up to 300 miles one way for screening or treatment. Some patients may be too ill, either from their disease or from treatment side effects, to make trips of this duration. Those with compromised immune systems utilize at their peril the van services a small number of communities have established. In some areas, volunteer transportation assistance programs help people reach treatment, but these too are limited.

Native Americans, particularly those living on reservations, frequently drive long distances to an IHS hospital or clinic and wait all day to be seen, only to be turned away and told to return the next day. This situation is so discouraging that some people avoid seeking care for symptoms until they require emergency care. According to the director of a screening program for Native American women, only 11 mammography machines are available to serve all 566 recognized Native American tribes. Even women with symptoms may wait months for a mammogram, and outreach and education efforts can create a demand for services that outstrips already limited capacity.

Most managed care enrollees and VA beneficiaries are required to obtain care only from providers with which
contracts have been negotiated. In many cases, rural patients must travel considerable distances—sometimes to a distant city or another state in the case of VA-approved treatment sites—to reach these providers. In some cases, they literally drive past equivalent treatment facilities near home on their way to the contracted provider.

Poor rural patients, particularly the elderly, often are not considered for clinical trials because of their physical isolation, transportation problems, and fixed income. In addition, clinical trial protocols are often rigid, not allowing monitoring and treatment schedules to be coordinated so as to minimize patient travel. This lack of flexibility results in additional long-distance travel for rural patients who do enroll in trials, discouraging participation.

Often, neither the patient nor his or her friends or family has a car that is sufficiently reliable to make long trips to the treatment facility; they also may not have the money for fuel and tolls. Many older Americans do not have anyone who can drive them to care and cannot afford to pay for transport. Since relatively few treatment centers are able to help patients with food or lodging expenses when they must remain near the treatment site for outpatient chemotherapy or radiation treatment, some patients end up sleeping in their cars, in homeless shelters, or on chairs in the hospital or clinic. In some cases, the time required to raise money to pay for transportation to treatment can mean the difference between life and death.

Transportation-related barriers and associated costs are difficult for rural and frontier area residents, but they are particularly onerous for patients in isolated regions like Alaska, Hawaii, the Virgin Islands, Puerto Rico, and the Pacific Territories. Often, the only way to reach screening or treatment of any kind is by air or boat. In Puerto Rico, for example, a poor cancer patient who misses the last ferry home following a chemotherapy treatment is faced with sleeping in the town square; most have no money and there is no other means of transportation. In Alaska, the cost to fly from a remote village to a medical facility providing colonoscopy may exceed the cost of the test itself. For those in areas that also have harsh winters, reaching cancer care may become impossible for weeks at a time when country and mountain roads become impassable due to snow. As speakers noted, such conditions can delay diagnosis of suspected cancer and seriously disrupt cancer treatment regimens.

“Because the cancer was so rare, there was no one in Rapid City who was qualified to do the surgery... so she was sent to Portland, Oregon... unfortunately, there wasn’t funds available to help her with the transportation and everything. So there was fund raisers that had to be done. So it took almost five months to get enough money together to get her up to the hospital and her husband to stay there with her. And after the surgery the doctor came in and said ‘I hate to tell you this; if you had been here four months ago, we could have gotten it all and you would be free from cancer. But it’s too late now and you have a year to live.’”

Dolly Randles, cancer survivor, South Dakota, describing her daughter’s cancer experience
Transportation also is an issue for many inner city residents who may not own cars or be able to afford bus or subway fares needed to reach screening or treatment facilities. Moreover, some urban patients are too sick to undertake a trip that may cover relatively few miles, but involves multiple bus transfers and considerable walking.

Technologies exist, or are emerging, that could help relieve some of the transportation problems faced by rural, frontier, geographically isolated, and inner city residents. Telemedicine has been tested in a number of rural settings and has shown considerable promise for bringing state-of-the-art diagnostic and other care to those at great distance from metropolitan cancer centers. Tumor boards, consultations, and continuing medical education are among the possibilities telemedicine may offer. Effective October 1, 2001, Medicare reimbursement for “telehealth” services will expand to include consultation, office visits, individual psychotherapy, and pharmacologic management delivered via specified telecommunications systems.

**What Barriers to Cancer Care Are Related to Information and Education?**

Lack of information or education, not knowing how to find or evaluate information, not believing or acting appropriately on available information, not knowing how to get needed care within the health system—all can constitute barriers to cancer care. Singly or in combination, they affect both health care providers and people who seek cancer information and care.

**Provider-Related Barriers Related to Information and Education**

Information and education barriers related to health care providers take several forms; these can seriously affect the care cancer patients receive and ultimately, affect disease outcome.

Lack of information about cancer and cancer care was the provider-related barrier most commonly described by those providing testimony at the regional meetings. The need to better educate primary care providers about cancer was emphasized repeatedly. Initial decisions about care can be a critical determinant of the patient’s outcome. Primary care providers, especially in rural and remote areas, often make these crucial referral and treatment choices.
Likewise, oncology professionals may not be aware of all available treatment options for a given type of cancer, and considerable confusion exists in the provider community about the strength of supporting evidence for and the validity of sometimes conflicting published screening and treatment guidelines. More broadly, there is confusion among providers, payers, and patients as to what constitutes quality care for cancer. Speakers suggested that more universal acceptance of guidelines and consensus statements could make it easier to practice consistent, high quality cancer care, and also could decrease the time needed to obtain treatment approvals and process claims. Clearer and more widely publicized guidelines also would help patients make more informed treatment decisions.

Staying abreast of the latest research in all aspects of cancer care is challenging for any practicing provider in the current health environment, but speakers indicated that clinicians in rural and other geographically isolated areas are at a particular disadvantage in this regard. Most continuing education programs and professional meetings are held in urban areas, and many rural providers also have no local network of peers with whom to exchange ideas and information. Nurses at many Native American health sites work entirely alone. It was suggested that information be provided in a format that can be quickly and easily absorbed. In addition, speakers encouraged the development of distance learning approaches for continuing medical education and computer linkages to help connect isolated providers.

The lack of reliable, up-to-date cancer care information, or a failure to change practice patterns based on new evidence, can have serious repercussions for patients. For example, the surgeon of a breast cancer survivor refused to discuss the option of lumpectomy, saying that all of his patients who had chosen breast conservation surgery had died within a year. Regional variations in the use of breast conserving treatment have been well documented, and have been attributed to a number of factors, including provider age, practice patterns, and bias. Other cancer patients reported that their providers expressed pessimism about their prognosis; in addition to robbing patients of hope, this attitude appeared to influence the treatment options that were offered. Many community providers are mistrustful of and do not understand clinical trials; in particular, few primary care providers have received any training about how clinical research is conducted. Providers also may unduly influence treatment choice through body language, to which anxious
patients are highly, if not consciously, attuned. Provider education is needed on clinical trials and on communicating information about treatment choices.

Lack of information also results in misdiagnoses and late diagnoses. For example, one testifier indicated that her physician performed a mastectomy for what he diagnosed as aggressive breast cancer; in fact, it was a spindle cell carcinoma that did not require such drastic surgery. Late diagnosis was particularly common among younger patients whose physicians dismissed or misattributed their symptoms to other causes. Several of the breast cancer survivors who testified, even those with symptoms and/or family histories of cancer, were told they were too young to have the disease; the same was true of younger patients with colon cancer. When these patients asked for screening or other tests, they often were denied because they did not fit screening guidelines or the typical patient profile. A woman with throat cancer testified, “I went from doctor to doctor, and I was told I was overreacting, that I needed professional counseling, that I wanted attention...and even worse, I was told I was a drug addict, because I had to keep working to support my family and I was asking for something to treat the [severe ear and jaw] pain.”

Speakers further suggested that providers are more likely to miss signs of cancer in patients with other chronic illnesses (e.g., diabetes, hypertension). They also may miss major health problems like depression in cancer patients, a problem recently documented in the literature. This problem may stem in part from the limited time physicians usually have to spend with each patient, since only the most urgent or newest presenting health issue may be assessed. Lack of provider knowledge also may contribute to this problem.

Many oncologists are not knowledgeable about hospice and instead are likely to refer terminal patients to intensive care, where end of life issues tend not to be addressed. In addition, many physicians are uncomfortable with and unskilled at discussing end of life care. Hospice providers indicated that referral to intensive care rather than hospice sometimes reflects an unwillingness on the part of the physician, the family, or both, to give up aggressive treatment of terminal patients.

Another problem described by speakers is a lack of provider information or education about cultural differences that affect the way patients need to receive information and how they perceive disease and treatment. This was distinguished from instances of bias, and from a more common lack of
communication skills and compassion ("bedside manner") that seemed to be at the root of unnecessarily blunt communications about diagnosis and prognosis.

Finally, an information problem cited by many providers involves the serious deficiency of existing medical data systems, including patient monitoring and tracking systems to help improve patient care and ensure that no one “falls through the cracks.” One physician observed that, “Medicine currently is less computerized than Wal-Mart.” Community providers related instances in which patients were referred for testing or treatment but did not receive it due to lack of transportation, lack of money, fear, or other reasons. With no monitoring or reminder systems in place, the providers sometimes did not realize that the additional care was not received until the patient returned months later with more advanced disease. One provider noted that whereas reminder cards and telephone calls for checkups and other appointments are commonly used by the dental profession—and even in nonmedical settings such as auto maintenance—to help ensure continuity of care and compliance with recommended treatment, such systems seldom are employed in cancer screening and cancer care programs.

Information and Education Barriers Faced by Patients and the Public

The lack of accurate cancer-related information that is readily available, understandable, clear, and delivered in a sensitive and culturally acceptable manner is a major contributor to the inability of patients and the public to obtain the most appropriate cancer prevention, treatment, and supportive care.

Cancer is perhaps the most feared of diseases. Few people have not been touched by cancer, seeing family members, friends, or co-workers succumb to some form of the disease. These experiences can be galvanizing lessons, particularly when they are not balanced by the experience of seeing people who not only survive, but live full and productive lives after having cancer. Fear of cancer can cause people to reject cancer prevention messages, avoid cancer screening, delay or avoid treatment if diagnosed, and have difficulty understanding and choosing among treatment options or following treatment regimens.

Across many populations and cultures, fear related to cancer takes the form of fatalism, fear of treatment and its costs, fear of pain and disfigurement, and fear of rejection by one’s partner, family, or community. In addition, myths
related to cancer persist in many populations, including beliefs that cancer is contagious, that surgery and exposing cancer to air causes it to spread, and that mammograms cause cancer. Overcoming these cancer-related fears and myths is a major challenge to those who provide cancer education and cancer services.

For some Americans, cancer-related information is simply unavailable. In remote regions and high poverty areas, thousands lack telephones. Some remote areas even lack radio or television reception. Library access is limited in many rural areas as well as some inner city areas. Few of the poor own computers or have access to the Internet. Many lower-income people without health insurance do not go to the doctor or to other sites where cancer information is most commonly found. Cancer educators and cancer control providers must learn where and how to reach such hard-to-reach populations. For example, it was suggested that cancer information should be made available at social welfare departments and other organizations that provide financial, housing, food, and other assistance. Health fairs, churches, and established community events have proven effective for reaching underserved populations. Some communities are making Internet information more accessible to populations that are unlikely to have such access at home or at work by having open computer hours at community health centers and holding free computer lessons for seniors.

For other Americans, cancer information may be readily available but entirely unusable due to literacy or language issues, or it may be presented in ways that are culturally unacceptable. Low literacy appears to be an underappreciated barrier to cancer information and appropriate cancer care. In a few states, as much as a third of the population cannot read. Low literacy seriously limits the ability to locate and comprehend information about cancer risk, prevention, early detection, or treatment options. Patients who cannot read may have difficulty following medication regimens or other physician instructions, and literacy level has been shown to be a factor in late stage diagnosis of cancer. 

Print materials available from government and voluntary organizations, health care organizations, and in newspapers and magazines are typically written at high school reading levels or above. Informed consent forms likewise are written at too high a reading level for many; these forms are intimidating even for many well-educated patients. Speakers reported some attempts to shorten and simplify both educational materials and consent forms for low literacy populations.

“My first experience with [a] cancer patient was a Cambodian older man who [had] stomach cancer. And he was...very lonely in his pain. And his family is a large family but they never went to help him...and so one day I asked the family, ‘Why don’t you do anything for him? Why don’t you touch him? Why don’t you talk to him? He’s going to die pretty soon.’ And they cried and they said they want [to] so much but they had to stand behind a glass door, they’re afraid that he will give them his cancer.”

Yung Kral, program director, Georgia

“There are people who are so afraid of the treatments...that they decide to leave it in God’s hands. I, myself, am a very strong believer, but I also believe that God has given us the medical establishment of which you must take advantage. Let us teach them that it’s okay to go to the doctor with God.”

Mavis A. Alleyne, cervical cancer survivor, New Mexico
Language barriers are a growing challenge to effective communication about cancer as the current wave of immigration continues. In Hennepin County, Minnesota, for example, the county hospital emergency department has used 91 languages to communicate with patients; in Los Angeles County, California, more than 120 languages are spoken. With so many languages (and distinct dialects of some languages) in use, hospitals and other health care providers are finding it extremely difficult to provide adequate interpreter services. In some cases, translation is provided by non-medical hospital workers or other strangers. Neighbors, friends, and even the children of patients are called upon to translate, which can be an embarrassing and awkward situation for all involved and may cause the patient to withhold important information. Moreover, the physician or other health care provider may not be able to assess whether information is being translated accurately. Written materials in numerous languages are costly and time consuming to produce. Speakers emphasized that many health-related materials translated from English versions are inadvertently translated either in the wrong dialect or at too high a literacy level. Some recent immigrants are not literate either in English or in their native language; for some population groups, picture books are needed to convey essential cancer information. Translation to certain languages is particularly challenging: in languages spoken by certain Alaska Native populations, there is no word for cancer, and in the native language spoken in the Northern Mariana Islands, there is no translation for “prostate cancer.” Disabled populations also may have special information needs; for example, the visually impaired may need large print or audio materials, and the hearing impaired may require sign language interpreters. Some local programs have developed tailored communication materials for specific populations. But with the exception of a small NCI-funded program to facilitate distribution of educational materials targeting Native American populations, there appears to be no established mechanism for collecting and disseminating such materials to others who may be able to use or adapt them for other population groups.

In addition to being understandable, clear, and in the right language, information must be presented in culturally sensitive and appropriate ways. The myriad cultures of the nation complicate attempts to provide cancer information, but cultural considerations must be incorporated into communication efforts. When the positive aspects of culture are...
woven into the design and presentation of cancer information, the message is far more likely to be accepted. A program director described how her staff helped to make cervical cancer screening relevant for a group of Native American women: “...they thought ‘cervix’ is a really strange word... [we said] think of it like this: ‘it’s the doorway to the world for your babies and grandchildren.’ And all of a sudden Pap smears became important to these women.”

Regardless of their educational level, most people need help to evaluate and prioritize information about cancer risk, prevention, and all aspects of cancer care. Just as too little information can lead to poor treatment choices, too much information also can lead to less than optimal care. Cancer information is now available from multiple sources, including print media (e.g., newspapers and magazines, pamphlets and other materials from research and advocacy organizations and health centers), television and radio, and the Internet. However, this material is presented at varying levels of detail, is of highly variable quality, and often contains conflicting recommendations. The Internet in particular is rife with persuasive information about untested alternative treatments. Information that is intended to be helpful can have unintended effects. For example, confusion about conflicting screening guidelines causes some people to avoid screening entirely. Inconsistent messages about cancer prevention discourage beneficial behavior changes. Media coverage of clinical trials that emphasizes problems rather than successes can contribute to fear of trials and discourage participation.

People newly diagnosed with cancer also are likely to be bombarded with information from family, friends, and health care providers. The physician’s role is changing from the principal provider of information to an interpreter of information. Many patients now arrive at the doctor’s office with information obtained from the Internet or other sources. Physicians and patients indicated that some physicians find this threatening, while others welcome the active involvement of patients in their care. Yet even with assistance, patients who choose to be involved in decisions regarding their care may themselves making life-altering, and even life or death choices with little confidence that they have received the best advice. In large measure, this is due to the fact that for the majority of cancers, the most effective care is not known.

A Massachusetts sarcoma survivor who faced the dilemma of inconsistent recommendations from doctors who rendered second opinions about her care, observed that physicians’ recommendations are subject to the individual provider’s
judgment based on the information available; patients ultimately have to go with their “gut feeling” in making treatment choices. At the other end of the continuum, it also was noted that in the Vietnamese medical system, doctors customarily do not explain to the patient why he or she is sick, or with what disease. They simply dispense medication and send the patient home. It has become part of the culture to not know the cause or to expect information about one’s illness.

Customary patterns of seeking health-related information can be a significant barrier to reaching people with accurate cancer information. For example, men of many cultures are uncomfortable discussing personal health issues, and so are unlikely to talk with each other about health problems or seek information in ways or in places that might disclose such problems. Therefore, as community program planners indicated, they are less likely than women to accept cancer education or screening efforts at the workplace. Targeting spouses and significant others to transmit colon and prostate cancer screening information and motivate men to obtain screening has proven effective in improving screening rates among at-risk men.

Seeking information through written material is a largely American and Western European cultural practice. Some cultures instead have a tradition of seeking health information from a trusted person in the community; however, such persons may not be knowledgeable about cancer. Several speakers described local programs designed to address this issue in which community members, usually women, are trained to provide cancer education, outreach, and support to peers. Variously known as community health representatives, lay health educators, promotoras, church health aides, or by other titles, these community members reach their neighbors—often through one-to-one communication or events held in homes or churches—with accurate and culturally acceptable cancer prevention and screening information.

Going beyond the need for information alone, however, patients and providers alike emphasized the critical importance of providing assistance in navigating the complex and fragmented cancer care system. The Panel heard testimony about several programs in which patients are assisted by individuals (e.g., patient navigators, ambassadors, patient advocates, case managers, patient liaisons, social workers) from entry into the screening process through definitive diagnosis and treatment of a detected abnormality and follow-up care. These navigators often help patients find out...
about financial and medical assistance for which they may qualify, help complete the required documentation, advocate for the patient with payer agencies, arrange transportation and child care, accompany patients to appointments, help obtain referrals and treatment approvals, find appropriate support groups, and provide emotional support to the patient and family. Importantly, the fiscal support of the programs described varied from small, purely volunteer efforts by grassroots organizations and churches, to grant-supported activities, to hospital system-wide programs funded by endowments. While the need for such programs for insured and uninsured patients at all educational and socioeconomic levels was unquestioned, the tenuous stability of these often fledgling programs was also underscored. An administrator from an endowment-supported hospital program that provides a case manager to every cancer patient upon entry into the system noted that due to reimbursement reductions, the hospital system has operated at a deficit for the past three years and the future of the program is now uncertain.

Information Barriers to Effective Cancer Control Planning and Implementation

Cancer surveillance is grossly underfunded in many areas of the country, and numerous presenters described protracted—and sometimes failed—efforts to secure state funding for these activities. They noted with frustration that funds for cancer, other health issues, or other non-health activities too often are based on institutional and political preferences or misperceptions of need rather than on data.

Without adequate information on the extent and nature of the cancer burden, states cannot identify high risk groups, focus their planning efforts, develop targeted prevention and cancer control efforts, or evaluate their success. Nonetheless, many states are pressing ahead with developing cancer control programs despite inadequate data on the cancer burden borne by populations within the state and lacking fundamental information on the care available and community needs. Some states, particularly those supported by NCI's Surveillance, Epidemiology, and End Results (SEER) program or the CDC's equivalent cancer registry program, have population-based registries. Other state cancer registries are not population-based, since only hospitals may be required to report. Even this hospital data may be less than complete; speakers noted that with many hospitals struggling just to stay open, cancer data reporting is not a priority. Moreover, a speaker pointed out that in

“...The [tumor] on my neck had grown so large that I didn’t know what to do... I was embarrassed because I didn’t have insurance. I was overwhelmed because I wasn’t getting anywhere [trying to find information and a source of treatment]...I wanted to give up...there’s people out there like myself in the crack or in the corner, whatever you want to call it, that don’t know where to turn.”

Albert Calloway, head and neck cancer survivor, Pennsylvania
independent-minded mountain states, it is difficult to mandate anything, including data reporting.

A positive development has been the recent expansion of SEER coverage through contracts to four additional states. This expansion effort, jointly funded by NCI and CDC, nearly doubles coverage of the U.S. population to over 65 million people and substantially enhances data collection on populations that bear a disproportionate cancer burden and about whom limited data currently exist. In addition, the recently developed state and county cancer mortality maps available on the Internet (http://cancer.gov/atlas) now provide state and local planners with data at a level of detail not previously available. However, many presenters emphasized the importance of detailed incidence, treatment, and survival data on local populations for guiding cancer control planning and implementation efforts, and data at this level remain sorely lacking in most states.

The lack of data for effective cancer control planning leaves most states and communities with a difficult dilemma. They recognize the importance both of devoting greater resources and effort to surveillance activities and of allocating scarce screening and treatment resources to data-driven interventions. Yet data on cancer patterns and trends may take years to develop, and surveillance activities should not supplant action to address readily apparent cancer screening, diagnosis, treatment, and support problems.

The Impact of Culture and Bias on Cancer Care

The population of the United States is becoming more diverse with each passing day. Elements of diversity (i.e., individual and population characteristics) include ethnicity, race, culture, religion, nation of origin, native language, literacy level and educational attainment, income level, gender, and age. In many important respects, these differences among people enrich the collective life of the nation, and the country has made strides in fostering a positive appreciation of and reducing bias based on diversity, particularly in the areas of employment, housing, education, and sport. Yet as a Nation, we still have far to go to understand and be fully respectful of individual and population differences.

Combinations of characteristics attributed to individuals or populations often are spoken of broadly as culture, including
characteristics attributed to perceived race. Culture also can be thought of as largely implicit frameworks of shared ideas, beliefs, social expectations, or habits of mind—one’s mental software.53 These frameworks, and our ideas about who and what we are, develop from our social environment, which in turn is influenced by history and chance. Importantly, these frameworks (or world views, or mindsets) exist for the most part below the surface of the individual’s conscious mind. People are seldom called upon to become conscious of or re-examine them unless confronted with mindsets that are substantially different, or when it is important to understand the mindset of another to achieve an objective.54

Thus, people bring their cultural mindsets with them to all of the experiences of their lives, including seeking cancer information and cancer care, or providing such information and care. Aspects of individual or group culture may contribute positively to these pursuits. However, to the extent these mindsets cause people to make health-related decisions not in their self-interest or in the interest of those who could be harmed by their decisions, the effect of culture is bias with negative outcomes.

Issues of Culture and Bias Originating With Health Care Providers

The cultural mindset of the health care provider has the potential and has been demonstrated to affect behavior and decision-making not related to an individual patient’s medical condition. As noted earlier in this document, to date; such disparities in treatment and disease outcome have been studied most intensively as they relate to care provided to African Americans compared with white Americans. Similar disparities also have been documented in the care of Latino populations, the elderly, women, and other population groups.

Health care providers have a special responsibility to be aware, in the immediacy of their contact with patients, of the potential negative impact of failing to understand and accommodate cultural differences—both their own and those of the patient. This responsibility extends to hospital and health plan administrators and other non-medical personnel who make decisions affecting patients’ access to information and care. This is especially true because of the power and authority many patients confer upon or accept from health care providers, particularly physicians.

Provider bias may be overt, but more often it appears to be both unintended and transparent to the health care provider,
agency, or institution. Speakers from the gay community testified to pervasive and often overt provider bias against gay and lesbian patients that causes these patients to avoid screening and care, to avoid disclosing their sexual orientation to providers, and to have few support services available to them. As a result, speakers maintained that lesbian cancer patients in particular are at greater risk of late diagnosis and poorer outcome. In addition, hospital policies often exclude the partners of gay and lesbian cancer patients from family discussions of or participation in the patient’s care.

Patients with disabilities may be denied screening or other cancer-related care because providers focus only on health problems related to the disability. Speakers suggested that this is particularly true for patients with mental, mobility, and spastic disorders.

In an instance of apparently unintended bias, an African American prostate cancer patient who held degrees in physics and mathematics, recounted that he was not told his Gleason score (a measure of tumor aggressiveness expressed as a number on a scale of two to ten) by his white physician because the doctor assumed he would not understand it.

Bias at the provider and institutional levels may also occur when assistance, referrals, treatment, and other services are offered more readily to more educated or white patients compared with less educated or minority patients, even when they have equivalent resources, or lack of them.

Speakers also suggested that some provider behaviors may reflect a lack of understanding and respect for cultural practices that are key to the effective treatment of specific patients. For example, physicians may refuse to allow patients or their families to accommodate traditional practices such as prayers or ceremonies in the hospital setting, even though doing so would not interfere with treatment and would be comforting to the patient. Speakers indicated that under these conditions, some patients reject further treatment.

In other cases, provider behaviors that impede the care process may stem from a simple lack of education about other cultures and their customs. To illustrate, a speaker explained that in many Native American cultures, people are always addressed by their first names regardless of station or situation. A non-Native health care provider who believes he or she is showing respect by addressing a patient more formally (e.g., Mr. Jones) is likely to be immediately dismissed as someone who does not understand the patient’s culture and world view.
Issues of Culture and Bias Originating With Patients and the Public

Culture and bias are not just issues for health care providers. As the paragraphs above suggest, cultural values and beliefs affect how—or even if—people approach and interact with the health care system and with individual providers. They also affect the ways in which people perceive illness, how they develop and act on medical and caregiver preferences including folk healing methods, how they explain and tolerate pain, and what they perceive to be quality care. Diverse cultural values and beliefs therefore can be a barrier to the most effective information and care if they cause people to make choices that result in less favorable medical and/or quality of life outcomes.

Fatalism about cancer pervades many cultural groups and affects the extent to which people are receptive to information about prevention and early detection, and their willingness to seek cancer care. Speakers noted that fatalism about cancer is widespread among Native Americans since people living on reservations with poor health care access have long observed that virtually everyone diagnosed with cancer dies from it. This view has engendered a disinterest in screening that educators about the benefits of early detection are challenged to change. Fatalism about cancer also is common among African American, Latino, and some Asian populations who believe that if one gets cancer it is God’s plan, a punishment from God, or the result of bad karma. Similarly, representatives from Appalachian states indicated that fatalism about cancer is common among this population, but rather than being seen as a punishment, the prevailing view is that everyone has to die from something, so it is pointless to seek screening or treatment.

The grinding circumstances and resultant culture of poverty profoundly affect the information and care seeking behaviors of the poor. Crucial daily activities such as finding and keeping employment, providing food and shelter, meeting transportation and child care needs, and remaining safe in sometimes hazardous environments leave little time for preventive health care, exercise, or other health promoting behaviors. In addition, since a great many of the poor are uninsured, few adults receive regular medical check-ups and are unlikely to take time off from work to seek care except for medical emergencies. These conditions, and the culture that accompanies them, are a major factor in the late diagnosis and limited survival of many poor cancer patients.

“Around the same time there was a baseball player that was diagnosed with [non-Hodgkin’s lymphoma], too. [Almost] the next day he was operated on...I still haven’t gotten any answer of every time that I ask how come it took four months for them to start treating me. So I’ve got to think that it was probably because I was not famous or maybe because I was Spanish...four months after they give you five years at the most. So why do they wait so long?”

Aide Montoya, non-Hodgkin’s lymphoma survivor, Maryland

“Poverty was what was on the minds of the people, and daily survival: how am I going to feed my kids today? How am I going to get out to the house in a four-wheel drive and back into town? Or how am I going to go to work and get my kids to day care? All of those daily survival things were a priority, not being screened for a disease that wasn’t even bothering them—it was unthinkable to do that.”

Delores Fallsdown-Geiger, outreach educator, Montana
Speakers also pointed out that rural residence and agricultural lifestyle comprise a distinct culture in which it is rare to seek medical care unless one is in significant pain. Rural pride dictates that medical services are not sought out unless one can pay the bill; since many farmers and other rural residents lack health insurance, care is likely to be deferred. People in small towns and farm areas also place a high value on privacy, and may avoid seeking care if it will mean disclosing one’s income or health problem. Some rural residents will only accept care if they can obtain it in another town where they are not known and have a means of transportation to get there. Speakers from rural states attested to the rigor of farm life, in which taking time to go to the doctor can mean postponing tasks whose timing is critical to a successful harvest. Women, it was noted, are essential to the day-to-day operations of the farm, and are unlikely to interrupt farm routines to seek health care even when they have symptoms. A similar tendency to minimize or hide their own health needs was noted among Asian populations. For example, a study comparing treatment choice for early stage breast cancer between Chinese, Japanese, and Anglo-American women found that the Asian American women were less likely to choose breast conservation (rather than mastectomy) because it would inconvenience others to take them to their radiation therapy or chemotherapy and prevent them from most rapidly resuming their family responsibilities. In addition, Asian women in the study tended not to choose oral chemotherapy not fully covered by insurance, since doing so would mean taking money out of the family’s savings, an expense deemed inappropriate unless it offered a clear survival advantage. This cultural view also was expressed movingly by a speaker from Hawaii, who recounted discovering by accident her mother’s ulcerated breast cancer that her mother had kept secret because she did not want to leave her children with bills. Her mother died shortly thereafter. Her daughter, also a cancer patient, indicated that if she did not have health insurance, she would do the same thing.

Provider-patient relationships built on familiarity and trust are crucial to effective education and medical care for some populations. Speakers indicated that trust relationships with providers are particularly important to Native Americans, yet patients at IHS facilities seldom see the same doctor. The same is true in other health facilities that tend to be staffed by temporary, rotating medical staff. Many of these facilities serve large numbers of patients who are new immigrants from cultures that also place a high value on a personal relationship with the provider.
For minority populations, issues of trust extend more broadly to a distrust of the health care system as a whole, which in many respects reflects a fear of mistreatment by people who have power, knowledge, and influence. African Americans in particular bear an enduring distrust of the health system, due in part to the infamous Tuskegee syphilis experiments and to a long history of substandard and disrespectful treatment that is shared by Latino and Native American populations. These experiences reinforce the fear of cancer treatment, including clinical trials, and are a significant barrier to care.

In some cases, resistance to entering the health care system and difficulty navigating it, is undergirded by cultural traditions concerning appropriate personal interactions. For example, Latino culture discourages challenging authority figures; among Native Americans, it is considered rude to be assertive or aggressive. Therefore patients from these cultures are less likely to be strong self-advocates.

Speakers testified to the critical need for health care providers from minority and underserved populations. Patients from these populations express a strong preference to be seen by providers from their same sociocultural group, but such providers are unavailable in most areas. As one speaker pointed out, there are only two Native American oncologists in the United States. In addition, more female providers are needed, since in Latino, Moslem, and Vietnamese cultures, for example, it is forbidden for a woman to be seen or touched by a man other than her husband. Thus, in Latino and other cultures in which men are the decision makers, women may be prohibited from going to male physicians for gynecologic examinations. Further, female providers are needed to help overcome issues of modesty common among older African American, Latina, and other women.

Presenters also described the special difficulty of reaching the Amish population with cancer education or cancer care. In this highly private and self-secluded culture, women are relatively uneducated and seldom receive cancer screening. These cultural barriers are not well understood.

In some Asian and other cultures, and particularly among newer immigrants from these cultures, it still is considered undesirable or inappropriate to disclose life-threatening illness to the patient because it may cause the patient to lose hope and die sooner. In strongly patriarchal cultures, the elder male family member typically makes all important family decisions, including those concerning health care. Though such practices often change with increasing acculturation, they may deprive
the person with cancer of the chance to achieve closure in personal relationships or otherwise prepare for death and can cause significant stress for younger-generation family members who do not agree with traditional practices.

Until relatively recently, discussing cancer was taboo in mainstream America. People spoke in hushed tones about “the big C” and cancer patients typically were said to have died “after a long illness.” As many speakers testified, prohibitions on discussing or disclosing cancer remain strong in many cultures. For example, many Native Americans believe that speaking aloud about cancer will bring it upon the reservation, and it is common for patients to keep their disease secret even from family members. Latina and African American women may hide their cancer because they fear rejection by their partners. In some Asian cultures, there is a strong fear of being shunned by both family and community. In relatively closed rural, Asian, and other communities that value self-reliance and privacy, there is a belief that emotional support should come from within the family, and that outside support is not necessary. If the culture also dictates that cancer is to be hidden even from family members, the person with cancer may die not only without treatment, but without the comfort and solace of family support. Taken together, these varied cultural beliefs create a “code of silence” that prohibits discussion of cancer, the result of which is delayed care or lack of care, needless suffering, personal isolation, and in many cases, preventable death.

**Conclusion**

The Panel is acutely aware that most of the issues and problems described in this document are not being expressed for the first time. Indeed, the very fact that these problems—faced by real people with cancer every day, in every corner of the nation—remain so prevalent makes concrete, achievable action to resolve them that much more urgent. We have heard these voices of our broken health care system—now we must answer.
Currently, 44 million people in the United States are uninsured, and at least 31 million of the non-elderly are estimated to be underinsured for a catastrophic illness such as cancer. Ample evidence indicates that even those with comprehensive health coverage may not receive the most appropriate care. Access to appropriate cancer care is the crucial fundamental step needed to relieve the desperate physical suffering, financial devastation, and loss of dignity so many people endure when cancer is diagnosed. If we lack the political will to craft and implement a national plan to address this unacceptable situation, then incremental steps must be taken to quickly remedy health care financing and delivery system elements that result in so much of the unnecessary distress now experienced by cancer patients and survivors and their families.

Continued research on the quality and equity of cancer care, outcome disparities, and related health economics and system issues is essential to guide transformation of the health care system in the coming years to better serve the public. But the people with cancer today, and their families, cannot wait for this distant relief. Therefore, the President’s Cancer Panel recommends:

**Immediate Action Steps**

1. Provide immediate medical coverage for the uninsured—84 percent of whom are workers and their dependents—upon a diagnosis of cancer to help ensure that no person with this disease goes untreated.

2. Address health coverage issues that contribute substantially to the financial devastation of people underinsured for cancer care costs:
   - Provide reimbursement for anti-cancer agents, supportive medications (e.g., antiemetics, pain medications), and proven chemopreventive agents regardless of method of administration.
   - Within two years, public and private payers should reach consensus on and implement a standard health benefit package for cancer care. This benefit package should be based on the best available medical evidence and should be updated regularly to reflect advances in the standard of care. The reports and deliberations of the Institute of Medicine, other groups, and consumers should be used to inform this effort.

3. Address patient and public needs for cancer information and for assistance in accessing services:
   - Provide funding to help communities coordinate, promote, and support community-based programs, including patient navigator programs, that help people obtain cancer information, screening, treatment, and supportive services.
   - Recognize that the services of non-physician personnel who are trained to conduct cancer screening, and provide cancer education and case management in varied settings are an important component of cancer care that should be reimbursed.
4. Sustain cancer care in the community by providing consistent and realistic health care provider reimbursement across states, and between urban and rural locations within states, for the cost of chemotherapy drugs and their administration.

**Longer-Term Solutions**

1. Medicare, Medicaid, the Veterans Administration, the Department of Defense, the Indian Health Service, and other public payers should clarify the order of responsibility for payment for cancer care services when individual patients are eligible for benefits under more than one program. This information should be communicated promptly and clearly to those who provide cancer care services and assist patients in navigating the health care system. The existing Quality Interagency Coordination Task Force may provide a forum for accomplishing this important task.

2. Develop Federal policies to minimize bias in the provision of cancer care:
   - Raise awareness of unintended or overt bias through initial and continuing training of health care professionals at all levels, as well as administrators and others who make decisions affecting patient care.
   - Establish and implement systems for monitoring treatment equity. In addition, expand quality of care research to include issues of treatment equity.

3. Minimize disparities in the provision of cancer care by:
   - Educating primary care providers about cancer.
   - Educating all cancer care professionals about the nature and application of evidence-based medicine and about clinical trials.
   - Developing and disseminating better tools to assist health care providers in conveying information about cancer and about cancer care options.

4. Address the problems of temporary medical staffing and cultural incompatibility by establishing additional mechanisms to encourage more minorities and members of other underserved populations to enter cancer care professions. Provide incentives to encourage providers to practice in medically underserved areas.

5. Extend state-of-the-art cancer care to rural, frontier, and other underserved areas by expanding the use of telemedicine and providing a reimbursement system that facilitates expansion of telemedicine to geographically underserved areas.

6. Permit more flexible use of categorical funding where appropriate to enable states to fashion more rational and more comprehensive cancer control programs.
1. President’s Cancer Panel. Cancer Care Issues in the United States: Quality of Care, Quality of Life, National Cancer Institute, 1999.


21. Ibid.


32. Harold P. Freeman, M.D., Chairman of the President’s Cancer Panel, was also appointed Director, NCI Center to Reduce Cancer Health Disparities on September 24, 2000.

33. President’s Cancer Panel, *Cancer Care Issues in the United States: Quality of Care, Quality of Life*, op.cit.


44. Detailed summaries of the testimony and town hall proceedings from all seven regional meetings are available at: http://deainfo.nci.nih.gov/ADVISORY/pcp/minsmenu.htm

45. The Health Insurance Portability and Accountability Act of 1996 (P.L. 104-191) has improved the ability of some insured persons with health conditions to change jobs and retain health coverage for pre-existing conditions if individuals can document continuous “creditable coverage.” However, those with a break in coverage of 61 days or longer are subject to a waiting period of up to one year for coverage of pre-existing conditions. In addition, the Act has other significant limitations; most importantly, it does nothing to ensure the affordability of health insurance or to provide coverage to the uninsured.


Appendix A

2000-2001 President’s Cancer Panel Regional Meeting Locations and Participating States and Territories

Omaha, Nebraska, June 15–16, 2000
- Illinois
- Indiana
- Iowa
- Kansas
- Michigan
- Missouri
- Nebraska
- Ohio

Burlington, Vermont, September 14–15, 2000
- Connecticut
- Maine
- Massachusetts
- New Hampshire
- New York
- Rhode Island
- Vermont

Billings, Montana, October 12–13, 2000
- Idaho
- Minnesota
- Montana
- North Dakota
- South Dakota
- Wisconsin
- Wyoming

Nashville, Tennessee, November 16–17, 2000
- Alabama
- Florida
- Georgia
- Kentucky
- Mississippi
- North Carolina
- Puerto Rico
- South Carolina
- Tennessee
- United States
- Virgin Islands

Los Angeles, California, February 1–2, 2001
- Alaska
- California
- Hawaii
- Nevada
- Oregon
- U.S. Pacific Territories:
  - American Samoa
  - Guam
  - Marshall Islands
  - Northern Mariana Islands
- Washington

Albuquerque, New Mexico, March 8–9, 2001
- Arizona
- Arkansas
- Colorado
- Louisiana
- New Mexico
- Oklahoma
- Texas
- Utah

- Delaware
- District of Columbia
- Maryland
- New Jersey
- Pennsylvania
- Virginia
- West Virginia
Appendix B

Figure A: Bringing Cancer Care Advances to the Public: Bridging the Divide Between Research and Delivery

<table>
<thead>
<tr>
<th>The Research Enterprise</th>
<th>The Delivery Enterprise</th>
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<tbody>
<tr>
<td>Basic Research</td>
<td>Delivery</td>
</tr>
<tr>
<td>Discovering New Knowledge</td>
<td>Applying and Providing</td>
</tr>
<tr>
<td>Translational Research</td>
<td>What We Know to All</td>
</tr>
<tr>
<td>Turning Discoveries</td>
<td>Improved Cancer Care</td>
</tr>
<tr>
<td>Interventions and</td>
<td>for All</td>
</tr>
<tr>
<td>Technologies</td>
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<tr>
<td>Applied Research</td>
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<tr>
<td>Proving New Interventions and Technologies Work; Learning to Apply Them</td>
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</tbody>
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What Activities?

- Laboratory Research
- Epidemiology
- Basic Behavioral Research
- Preclinical Research
- Phase I/II Developmental Trials
- Cancer Control Research
- Phase III/IV Trials (Large Scale Efficacy)
- Demonstration Studies
- Population Studies
- Epidemiology
- Development of Practice Standards
- Health Services Research
- Outcomes Research
- Health Care Coverage/Universal Access
- Public Education/Awareness
- Medical Education
- Dissemination of Practice Standards

Who Is Responsible?

- Universities
- Industry
- Other Private and Voluntary Orgs.
- NIH and Other Federal Agencies
- Universities
- Industry
- Other Private and Voluntary Orgs.
- NIH and Other Federal Agencies
- Public and Private Insurers
- Other Health Care Providers
- Legislators and Policy Makers
- Industry/Employers
- DoD, VA, HCFA, Medicaid and Other Public Payers
- Private Insurers
- Health Care Providers
- Universities
- Media
- Other Health Care Providers
- Educators
- Individuals
- Individuals

Source: Reuben, S.H., 2000. Adapted from Cancer at a Crossroads, Figure 2, 1994.
The National Cancer Program: Assessing the Past, Charting the Future
President’s Cancer Panel, Report of the Chairman, 1999