Organized patient activism has become a prominent feature of the biomedical landscape in recent decades. Most groups are disease-focused. AIDS activists invented the model. The breast cancer movement refined it. Bonnie J. Addario wants to perfect it for lung cancer patients.

Patient activists have raised funds for research, increased access to experimental therapies, redesigned clinical trial protocols, and accelerated regulatory approvals for new drugs. With a little help from her friends, Bonnie Addario is currently breaking new ground – she’s trying to invent a new institutional framework for cancer research.
Addario on their son High School Bonnie Addario, Woodrow Wilson High School (SF, CA), 1965; wedding day; Olympian Oil With second left to right: President of Company in South San Francisco. The company permanent secretarial position at the Olympian of menial jobs: “I did temp work for Kelly Girl. At car in the aftermath, and was forced into a series after twelve years with the family in dire financial man.” Bonnie married and began raising a family. ents were first-generation immigrants. My brothers by saying: “If someone had told me this would be of fortune. When she tells her life story, she begins with me, I would never have believed it.” Bonnie graduated from high school in 1965. “College wasn’t an option for me,” she says. “My parents were first-generation immigrants. My brothers were expected to go on. I was supposed to find a man.” Bonnie married and began raising a family. The union produced three children, but ended after twelve years with the family in dire financial straits. The young homemaker lost her house and car in the aftermath, and was forced into a series of menial jobs: “I did temp work for Kelly Girl. At night, I cleaned banks with my kids.” Suddenly, fortune smiled. Bonnie landed a permanent secretarial position at the Olympian Oil Company in South San Francisco. The company provided a range of services to gas and oil producers and marketers. Bonnie worked for the President and Chairman of the Board, performed well, and was rewarded with regular promotions. She displayed a knack for management: “I never went to business school. I figured it out along the way.” In little more than a decade, she became a retail manager, a supply distributor, an oil trader, and, finally, President of the company in 1997. Along the way, she had remarried. Life was good. Bonnie Addario’s success was unlikely, but her determination carried her through. Her untimely cancer diagnosis was a second potentially ruinous disruption in her life, and another test of will. “I poured myself a glass of wine and opened the newspaper. My eyes went to a small article about a new cancer care partnership between the University of California, San Francisco [UCSF] and Sequoia Hospital, close to my home in Redwood City.” The operation was a success. The surgeon removed the top left lobe of her lung, and part of her esophagus. The tumor was excised, although one of Bonnie’s vocal cords was paralyzed, and two major arteries in her heart were damaged and had to be repaired with grafts. After the surgery, Bonnie developed vascular necrosis, a side effect of steroids prescribed to moderate weight loss. The flow of blood to her bones was impeded. The bones became brittle, and she fractured her femur and hip as a result. She underwent two major surgeries to repair the breaks, and then, a few months later, was treated for a pulmonary embolism. When it was all over, Addario’s prog- nosis was upgraded. The ordeal left her physically depleted, but mentally charged. “My recovery was long and arduous,” she says. “I had plenty of time to think.” Bonnie’s first instinct after her diagnosis had been to understand her condition and recruit expert medical counsel. Her search for information had been aggravating. Institutional support and educa- tion for lung cancer patients was virtually non- existent. The lack moved Addario to contemplate executive action in a new territory: “I knew I had to educate people—patients, physicians, and the public—about lung cancer.” The cancer that the NCI forgot Bonnie embarked on advanced studies in the biology, epidemiology, and politics of lung cancer. In conversation with other patients, she learned that the ‘standard of care’ is late-stage diagnosis, a dearth of treatment options, and poorly-coordinat- ed medical services. She also learned that research on lung cancer is relatively neglected. In 1971, President Richard Nixon declared war on cancer. He signed the National Cancer Act into law, funneled a storm surge of federal dollars to...
In 2009 the National Cancer Institute funded over $3 billion in cancer research. Below is a comparison of NCI funding for six prevalent cancers. A significant disparity can be seen between the incidence and mortality of lung cancer and the research funding it receives relative to other cancers.

**The Stigmatized Cancer: Disparity in NCI Funding**

![Diagram showing NCI Funding](image)

**Source:** National Cancer Institute

**New Cases**
- Lung
- Colon
- Breast
- Prostate
- Non-Hodgkin Lymphoma
- Melanoma

**Deaths**
- Lung
- Colon
- Breast
- Prostate
- Non-Hodgkin Lymphoma
- Melanoma

NCI’s research support budget is allotted to lung cancer projects. In 2001, the NCI’s Lung Cancer Progress Review Group, a panel of thirty expert clinicians, scientists, industry representatives, and patient advocates, reported that funding for studies of lung cancer was “far below the levels that characterize other common malignancies and far out of proportion to its major incidence and mortality.” That’s not true. A comparison with breast cancer funding is telling. In 2008, lung cancer killed 161,840 Americans. NCI expended $14,145 per lung cancer death: $14,145. Addario questions the dedicated to early detection, diagnosis, and treatment. Addario objects to NCI’s funding priorities because studies of causation, prevention, and outcomes will not improve deficiencies in diagnosis and treatment — areas of pressing need that are not similarly deemphasized in, for example, breast and environmental factors – atmospheric pollutants such as radon or the fine particulate matter found in smoke, for example – contribute to the formation of lung cancer. Eighty percent of those diagnosed with lung cancer quit smoking decades earlier or never smoked at all. Addario objects to NCI’s funding priorities because studies of causation, prevention, and outcomes will not improve deficiencies in diagnosis and treatment — areas of pressing need that are not similarly deemphasized in, for example, breast research uncovers further issues. Sixty percent of patients die with lung cancer within eight to ten months of diagnosis. Addario believes that NCI’s emphasis on prevention has to do with the attempt to make sense of the world, to reduce the complexity of experience and discern order, but it’s simultaneously an attempt to assign blame. If lung cancer is the result of an individual’s decision to smoke, does it deserve as much attention as other common killers – breast cancer, prostate cancer, or colon cancer? Bonnie wants to set the record straight: “The world still assumes, apparently, that only smokers get lung cancer. That’s not true.”

Bonnie was eager to transform the ways in which lung cancer is understood and treated, and skeptical that universities could serve as catalysts of change. She decided to create a separate organization with a threefold mission – to provide patient support and advocacy, to educate and dispel misconceptions about the disease, and to heighten public awareness of the relative neglect of lung cancer research. With assistance from Whitney and Jablons, Bonnie established the Bonnie J. Addario Lung Cancer Foundation (BJALCF) in March of 2006. A few months later, the foundation made a public debut with a fundraising gala for lung cancer research in San Francisco. The evening drew a large crowd and raised an amount that was impressive (although “not nearly enough”). Bonnie was launched on her mission to fix a broken system. She determined to improve the dismal statistics on lung cancer, one patient at a time. The BJALCF’s initial efforts were directed toward patient education and empowerment. The foundation’s first public awareness campaign urged patients to get CT imaging scans for early detection. The message (“CT Scan – think of it as a mammogram for your lungs”) was emblemized in glowing pink and green on the sides of buses and highway billboards nationwide. The BJALCF’s mission was refrocused, however, and began organizing a diverse resource network through which patients and families could access information, support, and services. “We wanted to help people find their own best care,” Bonnie explains. “We connected them to experts, caregivers, and service providers at multiple sites. We focused on teaching people about what was available.” The goal was to help patients become their own advocates – just as Whitney and Addario had, with positive results. “Information is powerful,” says Bonnie. “Informed patients live longer. They just do.”

Bonnie was eager to transform the ways in which lung cancer is understood and treated, and skeptical that universities could serve as catalysts of change. She decided to create a separate organization with a threefold mission – to provide patient support and advocacy, to educate and dispel misconceptions about the disease, and to heighten public awareness of the relative neglect of lung cancer research. With assistance from Whitney and Jablons, Bonnie established the Bonnie J. Addario Lung Cancer Foundation (BJALCF) in March of 2006. A few months later, the foundation made a public debut with a fundraising gala for lung cancer research in San Francisco. The evening drew a large crowd and raised an amount that was impressive (although “not nearly enough”). Bonnie was launched on her mission to fix a broken system. She determined to improve the dismal statistics on lung cancer, one patient at a time. The BJALCF’s initial efforts were directed toward patient education and empowerment. The foundation’s first public awareness campaign urged patients to get CT imaging scans for early detection. The message (“CT Scan – think of it as a mammogram for your lungs”) was emblemized in glowing pink and green on the sides of buses and highway billboards nationwide. The BJALCF’s mission was refrocused, however, and began organizing a diverse resource network through which patients and families could access information, support, and services. “We wanted to help people find their own best care,” Bonnie explains. “We connected them to experts, caregivers, and service providers at multiple sites. We focused on teaching people about what was available.” The goal was to help patients become their own advocates – just as Whitney and Addario had, with positive results. “Information is powerful,” says Bonnie. “Informed patients live longer. They just do.”

Bonnie was eager to transform the ways in which lung cancer is understood and treated, and skeptical that universities could serve as catalysts of change. She decided to create a separate organization with a threefold mission – to provide patient support and advocacy, to educate and dispel misconceptions about the disease, and to heighten public awareness of the relative neglect of lung cancer research. With assistance from Whitney and Jablons, Bonnie established the Bonnie J. Addario Lung Cancer Foundation (BJALCF) in March of 2006. A few months later, the foundation made a public debut with a fundraising gala for lung cancer research in San Francisco. The evening drew a large crowd and raised an amount that was impressive (although “not nearly enough”). Bonnie was launched on her mission to fix a broken system. She determined to improve the dismal statistics on lung cancer, one patient at a time. The BJALCF’s initial efforts were directed toward patient education and empowerment. The foundation’s first public awareness campaign urged patients to get CT imaging scans for early detection. The message (“CT Scan – think of it as a mammogram for your lungs”) was emblemized in glowing pink and green on the sides of buses and highway billboards nationwide. The BJALCF’s mission was refrocused, however, and began organizing a diverse resource network through which patients and families could access information, support, and services. “We wanted to help people find their own best care,” Bonnie explains. “We connected them to experts, caregivers, and service providers at multiple sites. We focused on teaching people about what was available.” The goal was to help patients become their own advocates – just as Whitney and Addario had, with positive results. “Information is powerful,” says Bonnie. “Informed patients live longer. They just do.”

Bonnie was eager to transform the ways in which lung cancer is understood and treated, and skeptical that universities could serve as catalysts of change. She decided to create a separate organization with a threefold mission – to provide patient support and advocacy, to educate and dispel misconceptions about the disease, and to heighten public awareness of the relative neglect of lung cancer research. With assistance from Whitney and Jablons, Bonnie established the Bonnie J. Addario Lung Cancer Foundation (BJALCF) in March of 2006. A few months later, the foundation made a public debut with a fundraising gala for lung cancer research in San Francisco. The evening drew a large crowd and raised an amount that was impressive (although “not nearly enough”). Bonnie was launched on her mission to fix a broken system. She determined to improve the dismal statistics on lung cancer, one patient at a time. The BJALCF’s initial efforts were directed toward patient education and empowerment. The foundation’s first public awareness campaign urged patients to get CT imaging scans for early detection. The message (“CT Scan – think of it as a mammogram for your lungs”) was emblemized in glowing pink and green on the sides of buses and highway billboards nationwide. The BJALCF’s mission was refrocused, however, and began organizing a diverse resource network through which patients and families could access information, support, and services. “We wanted to help people find their own best care,” Bonnie explains. “We connected them to experts, caregivers, and service providers at multiple sites. We focused on teaching people about what was available.” The goal was to help patients become their own advocates – just as Whitney and Addario had, with positive results. “Information is powerful,” says Bonnie. “Informed patients live longer. They just do.”
“Fourteen Years and Counting”: Wells Whitney on Iressa®

When Wells Whitney started taking the cancer drug Iressa® in 2001, he wasn’t sure what to expect. After his diagnosis in 1998, he combatted his cancer with the standard weapons: surgery and chemotherapy. A surgery removed a large main tumor, successive rounds of chemotherapy reduced the number of remaining nodules from nineteen small active growths to four dormant masses. He hoped Iressa® would help him sustain this heartening progress.

If chemotherapy is a sledgehammer and surgery a knife, Iressa® is a bow and arrow. Developed by British pharmaceutical company AstraZeneca for the treatment of advanced non-small cell lung cancer, the drug is designed to disrupt epidermal growth factor receptors (EGFR) on lung cancer cells, and block signals that promote tumor cell growth. Unlike chemotherapy, Iressa® targets a specific molecular pathway.

For Wells, it worked. He began taking Iressa® in 2001 as part of a compassionate use clinical trial, two years before the drug received accelerated FDA approval. Research showed that Iressa® shrank tumors in 10% of patients. Wells was a lucky ‘ten percent.’ “The drug transformed his fatal cancer into a chronic disease,” he says. “I still have cancer,” he says. “I’m not cured, but I’m dealing with it.”

In 2004, researchers at the Massachusetts General Hospital, Sloan-Kettering Cancer Center solved the selective response riddle. They showed that patients with a mutation on the EGFR gene tend to respond positively to Iressa®. The mutated EGFR receptors are ten times more responsive to the drug than normal receptors. The mutation is most common in Asians, women, and non-smokers. “According to the genetic profile of my tumor,” says Wells, “I’m much more like a non-smoking Japanese female than a Caucasian male.”

Bad news arrived in 2005. Further trials of Iressa® revealed that patients taking the drug had a median survival of 5.6 months compared to 5.1 months for those on the placebo. Citing a lack of clear efficacy in life extension, the FDA took Iressa® off the market and restricted access to current and previous users.

Fourteen years after his diagnosis, and eleven years since his first dose of Iressa®, Wells continues to walk among the living. He is part of a small group of patients that has access to the drug through a mail-order pharmacy. For new lung cancer patients, Iressa® is no longer an option, but Tarceva®, another EGFR-inhibitor developed by Genentech and OSI Pharmaceuticals, has been shown to extend survival, is approved by the FDA, and is available to patients.

“If money were no object…”

Patient empowerment was one piece of the puzzle. Addressing inequities in biomedical research was another. Bonnie saw that embedded attitudes and practices in the biomedical establishment needed to be perturbed and recast, and that the BJALCF needed to recruit thought leaders in lung cancer research to the cause. In order to make an impact on the conduct of laboratory and clinical research, the Foundation would need a bigger stage and the firm support of influential experts in science and medicine.

Toward these ends, the BJALCF organized a meeting of world leaders in lung cancer research and care. On November 9, 2007 the best and the brightest in cancer research gathered in Genentech Hall on UCSF’s new Mission Bay campus for the first annual BJALCF Lung Cancer Summit. Wells Whitney served as master of ceremonies. Attendees included representatives from the NIH, Genentech, CancerCare, an oncology patient support organization, and Changing Our World, a philanthropy and fundraising consulting group.

The keynote speaker for the two-day conference was Michael Milken, financier, philanthropist, cancer survivor, and founder of FasterCures, a self-described ‘action tank’ working to improve biomedical research. The all-star roster of physicians and cancer researchers on hand included Jablons, Dr. Mark G. Kris of the Memorial Sloan-Kettering Cancer Center, Dr. Fred Marcus of Sequoia Hospital and Dr. Harvey Pass of the NYU School of Medicine.

As the researchers rose in succession to talk about their research, Bonnie realized that ingrained conference habits are hard to break. The summit was struggling with a terminal illness of its own: ‘death by powerpoint.’ Addario intervened, drew a curtain on the long string of self-congratulatory progress reports, and refocused the meeting. “We don’t want to know what you’re doing,” she said. “We want to know what you would do to increase lung cancer survival rates?” Bonnie encouraged the group to momentarily forget the obstacles and imagine the possibilities.

To be continued…

Part II recounts how Bonnie J. Addario, in pursuit of improvements in lung cancer research and care, has pushed patient activism in new directions. She has formulated original ends for the BJALCF and devised innovative strategies for achieving them. Stay tuned.