2009 REPORT TO THE BRAIN TUMOUR COMMUNITY
During 2009 generous donations of money, time and talent brought hope to even more people affected by brain tumours across Canada.

Our donors continued to be generous and dedicated to supporting the efforts of Brain Tumour Foundation of Canada. As a result of a favourable outcome of revenues over expenditures, we were able to allocate a substantial portion of this excess to our growing focus on funding research grants, as well as to the development of an updated and more user-friendly website in 2010.

In this report back to you, our community, we extend our sincerest appreciation for the positive impact of your support. Together we will find the cause of and cure for brain tumours while improving the quality of life for those affected.

Thank you everyone,
**SUPPORT**  
*People Helping People*

In 2009 Brain Tumour Support Groups were implemented in Burlington, ON and initiated in Fredericton, NB. Due to the expansion of the program and increased awareness efforts, the total number of support group members in 2009 increased to 847, a 30 per cent increase from the 650 members in 2008.

“It is vital to keep on growing. The Foundation is an amazing group that does incredible things on a shoestring budget. The kind of support they offer is just remarkable, and it’s in the process of getting better.”

Jane Arnott, mother of 22-year-old son Ben, a brain tumour survivor

**FAMILY SUPPORT**  
*Respite, Education & Sharing*

Throughout 2009 the BrainWAVE Program – led by dedicated volunteers and parents – continued to grow with ongoing expansion in Lower Mainland British Columbia and increased enrollment in Southwestern Ontario.

“The program provides fun opportunities for children and teens to connect with others with similar experiences while giving parents the chance to learn, share information and network with other parents. Through the BrainWAVE Program, families are able to inspire each other with hope and support.

“As parents, BrainWAVE provides a great opportunity for Sarah to have a lot of fun with kids who are like her. Sometimes that’s important. Where do you find the new normal after brain tumour treatment? At BrainWAVE events it doesn’t really matter. The kids have fun and it lets parents deal with the special needs that we have around our children and their brain tumours. The information sharing is invaluable.”

Russell Smith, Sarah’s Dad

**HANDBOOKS**  
*Empowerment through Information*

During 2009, 6,915 adult and pediatric Patient Resource Handbooks were distributed free of charge across Canada and internationally to meet the needs of survivors, family members, caregivers and health care professionals.

“Many devour it, saying how great it is. Patients and families especially like the diagrams and glossary, because the world of brain tumours presents a whole new language.”

Marianne Lee, Neurosurgical Social Worker

“You are so overwhelmed when your child gets the diagnosis. The handbook gives you security and power. You feel so helpless. You can’t fix your child, you don’t understand the problem and you don’t understand the potential solutions. With the pediatric handbook I could read and then understand when I spoke with Sarah’s doctors”.

Russell Smith, Sarah’s Dad

**HELPING CHILDREN**  
*Making a Difference in Young Lives*

During 2009, 2,059 children storybooks, in English, French and Italian, about a little girl with a brain tumour and her journey through her treatment were delivered free of charge to families, children and schools across Canada and internationally.

“I love to share ‘A Friend in Hope’ with children. With each word I read, each picture I share, I can feel, see, and hear their joy, sadness, laughter, love. They are eager, open to sharing their feelings, asking questions, wanting to help. And then one child softly shares, ‘Miss Pia, my sister has a brain tumour.’

The storybooks make a difference in children’s lives.”

Pia DiBacco, Volunteer who introduced the children’s storybook to the Montreal school system and was the impetus and fundraiser for the translation of “A Friend in Hope” into the French version “Mon Amie Claire.”
EDUCATION
Information Day Conferences
Leading researchers and brain tumour experts were keynotes at the annual Halifax, London and Calgary Brain Tumour Information Days. Dr. Vivek Mehta, Neurosurgeon from the University of Alberta, Dr. Peter Dirks, Pediatric Neurosurgeon at The Hospital for Sick Children in Toronto and Dr. Brian Thiessen, Neuro-Oncologist at the BC Cancer Agency all shared their expertise along with other leading healthcare professionals.

Topics ranged from a look at the role of stem cells in brain tumour growth to advances in brain tumour treatment.

In 2009, 820 people from across Canada attended in a warm, comfortable and supportive atmosphere for survivors and their families. They were able to connect with one another, empower themselves with knowledge from experts in the field and ultimately find hope.

RESEARCH
Funding Hope
One of the highlights of 2009 was the August launching of the William Donald Nash Brain Tumour Research Fellowship, to advance the Brain Tumour Foundation of Canada’s vision to find the cause of and cure for brain tumours while improving the quality of life for those we serve. The fellowship will run for two years and will include funds to cover salary and lab expenses of $50,000 per year.

This Fellowship will encourage and enable young clinicians and scientists to enter the field of brain tumour research and will accelerate studies leading to a cure for brain tumours. The expectation is that Bill Nash’s generous gift will allow us to fund at least five fellowships over the next 10 years.

Six brain tumour research projects with grants totaling $146,714 were awarded in 2009. There were 34 applicants to the annual program and the six Canadian researchers will be conducting research projects focusing on causation, diagnosis and treatment. One grant was awarded specifically to investigate the role of genetic and environmental factors in the development of pediatric brain tumours.

ADVOCACY
Speaking Out for Better Healthcare
As a member of the Canadian Alliance of Brain Tumour Organizations (CABTO), Brain Tumour Foundation of Canada participates in CABTO’s efforts to advocate for improved patient care, increased government funding for brain tumour research in Canada, and increased public awareness. Two issues were of particular note in 2009:

1. The need for better data collection around the incidence of brain tumours in Canada. In 2009, a report was prepared through the Public Health Agency in cooperation with the Council of Cancer Registries. It outlines opportunities to improve the consistency and completeness of statistics about brain tumours and is being considered by the Council of Cancer Registries for implementation.

2. Equitable access to and coverage for cancer drug therapies in all provinces. During 2009, CABTO members continued to learn about the differences between provincial access and coverage for drug costs with the hope of creating positive change where possible.

In 2009, the traveling Hats Tribute collection grew to 140 hats, raising awareness about people affected by brain tumours while on display at conferences, open houses and various community events. Each hat tells an inspiring story of courage on a personal journey with a brain tumour.

"You can’t put a price on Information Days. I’ve learned so much from the excellent speakers. You meet people from all over and sharing experiences gives you a great sense of hope.”

Tom Hicks, brain tumour survivor

"There is always something new and more hopeful. It’s especially encouraging when your specific type of treatment is the topic of a presentation. Having a brain tumour and the associated treatments is not easy.”

Annele Robertson, brain tumour survivor

With high quality brain tumour tissue samples critical to advances in research, the Brain Tumour Tissue Bank funded entirely by Brain Tumour Foundation of Canada, collected and banked samples from 67 patients in 2009. These samples yielded 369 samples which were added to the library of nearly 1,500 samples available for researchers.

"Bill’s generosity has created a legacy of support for young people interested in brain tumour research and will help to move our joint mission forward, to find the cause, better treatments, and one day a cure for brain tumours.”

Dr. Joseph Megyesi, Neurosurgeon and Chair of the Board of Brain Tumour Foundation of Canada

"I am very grateful for research funding, that is what brings hope for people who are suffering.”

Annele Robertson
Kelly Northey

Volunteers
The Heart of our Services

Brain Tumour Foundation of Canada was proud to recognize four volunteers who embody the spirit of volunteerism with the presentation of the Volunteer of Distinction Awards. These volunteers represent the more than 700 people who generously donate their time to helping brain tumour patients.

The Volunteers of Distinction honoured in 2009 were: Claudine LePage of Moncton, New Brunswick; Crystal Tellett of Calgary, Alberta; Robin Bateman of London, Ontario and Melissa Martin of Windsor, Ontario.

"Volunteering gets me out of the house and makes me forget about my rheumatoid arthritis. I love the staff and the work that I do, helping get thank you letters and receipts to donors as quickly as possible. My volunteer role, now that I’ve learned the donor database system, makes me feel needed and productive."

Robin Bateman, Administrative Volunteer

"Seven years after I began volunteering for the Support Group, some of the same members are still here and it’s like a little family. I love seeing the progress people in the group have made and how everyone helps new members cope with a diagnosis. Just knowing that you made a little difference makes a huge difference to me, I just like giving back and it does feel great to know that you may have helped someone."

Claudine LePage, Moncton Support Group Volunteer Facilitator

Heritage Club members ensure support, education and research funding for the future through making a legacy gift in their wills or through insurance policies.

Joe and Sue Barnes
Leslie Carter
Peter and Fairda Chislett
Katie Dark
Carina Jacobson
Barbara Kennedy-Murray
Donna McKee

Susan Marshall
Grace Schenk
Michelle Thibodeaux
Shelly Thierman
Lois Toll
Valerie Wicks

"I’m sure the money will help with research and the other important work being done by the foundation. I’ve had a good life, in fact I’m still living a good life and I just want to help."

93-year-old Leslie Carter, who makes annual donations to the Foundation, in addition to a significant bequest in his will.

Kelly Society

Named for Kelly Northey, the 8-year-old who inspired the founding of Brain Tumour Foundation of Canada, this special society honours the generosity and multi-year commitment of its members.

Ken Arnott
Chantalle Butler
Ted Cooney
Pam & Rolando Del Maestro
Chaua Horner
Patricia Klein
Maryanne MacDonald
Heather Mastromattei
Susan Marshall

Joseph Megyesi
Melodie and Steve Northey
Phyllis Retty
Ben Seewald
Michelle and Sean Taggart
Beth Tyndall
Fred Wilder
Peter & Susan Yates

Donors are the Lifeblood of all the services and research funding provided by Brain Tumour Foundation of Canada. We salute you all for your generosity and your commitment to making a difference for those making the journey with a brain tumour.
SPECIAL MAJOR GIFTS

We salute those donors who have made major gifts to help in 2009.

- Estate of Andrea Dalton Baker
- The Crone Family
- Alison and Darrell Jones
- Joseph Megyesi
- Estate of Patricia Moore

LEGACY FUNDS

These special Legacy Funds continue to provide funding to chosen services thanks to the ongoing support of our legacy families.

- The David Bloom Legacy Fund
- The Kelly Northey Legacy Fund
- The Hannah Patterson Legacy Fund

CORPORATE AND FOUNDATION DONORS

Thanks to those corporations and foundations that help significantly through sponsorships or through special program funding.

- Deloitte & Touche Foundation Canada, Platinum Sponsor
- Schering-Plough Canada Inc., Gold Sponsor
- AstraZeneca Canada, Inc., Silver Sponsor
- IMRIS Inc., Silver Sponsor
- WestJet, Silver Sponsor
- Hoffmann-La Roche Ltd., Bronze Sponsor

In addition to corporate sponsorships, special gifts were received from the following corporations and foundations:

- A.M.A. Plastics Ltd.
- Great West Life Assurance Company
- HB Group Insurance Management Ltd.
- Marchand Fairchild Blais Financial
- North Waterloo Farmers Mutual
- Par Fore Marketing
- Vivid Creative Communications
- RBC Foundation
- Suncor Energy Foundation
- Sun Life Assurance Company of Canada
- The Plunkett Foundation
- The Valley Gourmet

MONTHLY GIVING CLUB

Our monthly donors provide important and sustainable support.

- Lewis Balsdon
- Tony Barton
- Faheem Bohkari
- Iona Bolger
- Diane Bradshaw
- Keith Butling
- Wendy Camp
- Courtney Channon
- Debbie Collins
- Claude Cote
- Irene Cranstone
- George Daniel
- Miriam Deveerett
- Isabella Di Cristofaro
- Marie Duff-Whicelchel
- James Dynes
- Justin Erdman
- Lary Ewanchuk
- Marianne Franklin
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- Muriel Grant
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- Brian Marriott
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- Joan Meddie
- Brian and Jo-Anne Montag
- Carolyn Naas
- Catherine Nemeth
- Marian Parker
- Everett Pope
- Farida Rasulbakhsh
- Nancy Rideout
- Carolyn Ross
- Ruth Russell
- Justin Shannon
- Patricia Sharpe
- Wendy Sidwell
- Fred Siemonsen
- Ralph Simons
- Erin Somerville
- Cara Stein
- Gwladys Tanner
- Graham Thompson
- Albert and Rosemary Toews

HOW WE REACHED MORE CANADIANS AFFECTED BY A BRAIN TUMOUR IN 2009

% OF SPENDING

- RESEARCH 30%
- SUPPORT SERVICES 29%
- INFORMATION SERVICES 19%
- EDUCATION & AWARENESS 22%
- FUNDRAISING 24.3%
- PROGRAMS 72.3%
- ADMINISTRATION 3.4%

REACHING CANADIANS

In 2009, our gross revenue totaled almost $2.23 million and our expenses totaled $1.95 million. With this positive position, we have been able to direct more dollars to support programs, and particularly research, thanks to the generosity of our donors.

SPECIAL EVENTS

Thank you to the thousands of people who supported the 5,527 participants in the 22 Spring Sprints in cities across Canada in 2009.

Thanks to the families, friends and co-workers of brain tumour patients who raised funds through the 97 Community Events held in their honour or memory.

“Just seeing so many people get together at the Spring Sprint was great. An aunt of ours had a friend in Iowa who contributed and there was our uncle Ted from Australia – it’s just amazing to see people from different areas come together to give money and offer their support.”

Jennifer Fagan, mother of 11-year-old Max, a brain tumour survivor
VISION
To find the cause of and cure for brain tumours while improving the quality of life for those we serve.

MISSION
To reach every Canadian affected by a brain tumour through support, education, information and research.

VALUES
These serve as our guidelines for our conduct and behaviour as we work toward achieving our vision:

- Hope
- Caring
- Integrity
- Accountability
- Collaboration
- Connecting the Brain Tumour Community

620 Colborne Street, Suite 301, London, ON N6B 3R9
519-642-7755 1-800-265-5106

Charitable Registration No. 118816339RR0001