Follow up Care for Breast Cancer Patients

Living Beyond Cancer

what happens now?
Your cancer treatments are now finished. You are joining a growing number of people living with cancer who are often called cancer survivors. Follow up care is a very important part of your cancer care.

This guide describes your recommended follow up care plan. It will help you learn how to manage your life after cancer treatment, understand what you can do to reduce your risk of the cancer coming back and explain the tests that are recommended for you. Your cancer doctor has also sent your health care providers (family doctor/nurse practitioner and, if needed, surgeon) a copy of your follow up care plan.

Your family members may also be at risk for breast cancer and should talk to their health care providers about their cancer risk and the screening that is right for them.

You and your health care providers are an important part of your follow up care. Your health care provider can contact your cancer doctor anytime. If needed, you will be referred back to your cancer doctor.

We hope this guide will be helpful. If you have any questions or concerns, please talk to your family doctor/nurse practitioner/surgeon or cancer patient navigator (if there is one in your area).

Please know that you are not alone as you move forward in your recovery.
What is Follow up Care?

The goals of follow up care are to:

- Provide support. Follow up care visits allow you to discuss how you are doing and any problems you have, such as dealing with the side effects of cancer treatment and fear of recurrence.

- Check for the return of cancer or a new cancer.

Follow up care for breast cancer involves:

- Regular visits with your health care provider. These visits are very important to assess and talk about how you are doing and your current health. The visit will include a breast exam, feeling for lymph nodes in your neck and armpits and a general check-up. Blood work is not usually suggested for patients with non metastatic breast cancer (cancer that has not spread beyond the breast).

- A mammogram every year until age 75. After 75 please discuss with your doctor to see if it is beneficial to continue with annual mammograms.

- You may want to perform self breast examinations. If you do, talk with your health care provider about the best way to do so.

Everyone in follow up care will not have exactly the same care because each follow up plan is based on your treatment and how it affected your body.

This plan is a guide and does not replace your health care provider’s advice.
Follow up Care after Breast Cancer<sup>1</sup>

<table>
<thead>
<tr>
<th>Follow up Care</th>
<th>Year 1</th>
<th>Year 2</th>
<th>Year 3</th>
<th>Years 4 and 5</th>
</tr>
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<tbody>
<tr>
<td>Health care provider visit</td>
<td>Every 4 months</td>
<td>Every 4 months</td>
<td>Every 4 months</td>
<td>Every 6 months</td>
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<td>Mammogram</td>
<td>Annually until age 75 and then at the discretion of your health care provider.</td>
<td>Annually until age 75 and then at the discretion of your health care provider.</td>
<td>Annually until age 75 and then at the discretion of your health care provider.</td>
<td>Annually until age 75 and then at the discretion of your health care provider.</td>
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After year 5, you should have annual visits with your health care provider.

<sup>1</sup> American Society of Clinical Oncology, 2006
Talk to Your Health Care Provider About

- **Birth control**: Birth control methods that use hormones (such as pills, patches, injections or NuvaRing) are not suggested for women who are breast cancer survivors. Your health care provider can suggest other effective birth control methods.

- **Post-menopausal hormone replacement therapy**: Hormone replacement therapy may not be recommended. If you have menopause symptoms, ask your health care provider what you can do.

- **Menopause caused by treatment**: Symptoms can include hot flashes, mood changes (such as depression), dry vagina, painful sexual intercourse or loss of sexual interest. If you have any of these symptoms ask your health care provider what you can do.

- **Risk of osteoporosis (bones that break easily)**: All post menopausal women and women who have received aromatase inhibitor therapy are at risk for osteoporosis. Your breast cancer treatment may also increase your risk. There are ways to decrease your risk of osteoporosis, such as healthy eating and calcium supplements. It is recommended that you take Vitamin D 1000 IU daily. Regular weight bearing exercise will help to prevent osteoporosis. If you do develop osteoporosis, your doctor may suggest treatment.

- **Limit or Avoid Alcohol**: Discuss risks associated with alcohol use with your health care provider. Drinking alcohol increases your risk for several types of cancer. Risk increases for some kinds of cancer with as little as one drink per day. The more you drink, the more your risk increases. The risk of cancer increases regardless of what type of alcohol you drink (beer, wine, cider, spirits or liqueur). If you do drink alcohol, follow the Canadian low risk drinking guidelines. If you do not drink alcohol, don’t start.

  Combining smoking with alcohol increases cancer risks even greater than the cancer risks that come from drinking alcohol or smoking on their own.

- **Stop Using Tobacco**: Quitting smoking or other tobacco use after a cancer diagnosis does make a difference. It can result in fewer treatment and cancer-related complications. It can make your cancer treatments more effective, improve survival and reduce the chance of a recurrence or a second cancer. It is never too late to quit tobacco. Help is available. If you want to quit, talk to your health care provider or call 811 to be connected to a tobacco cessation counselor who can offer assistance over the phone and refer you to a quit program in your area. There is no cost and no referral is required. Online resources are also available at: https://tobaccofree.novascotia.ca/

  *Tobacco includes smoked tobacco (cigarettes, pipes, cigars) and smokeless.

Continued on next page
Talk to Your Health Care Provider About (continued)

**Use of complementary therapies:** Many therapies such as massage therapy, reiki and therapeutic touch can be helpful. If you are thinking about using complementary therapies, please talk with your health care provider to make sure they will not interfere with cancer treatments.

**Inherited risk of breast cancer:** The majority of breast cancer (90%) is not inherited (passed down through the family). But, it is important to identify people who could be at risk of inherited breast cancer; their screening and management options are different from a person who developed breast cancer by chance. It is important to review your personal and family history of cancer with your health care provider to see if you should be referred for a genetic assessment. Because this decision is based on family history and your own personal medical history it is important to have accurate information about the cancer in your family. Tell your health care provider the age people were diagnosed with breast (one or both sides), ovarian and male breast cancer, and the age of any family members when diagnosed with cancer. Not everyone who is referred for genetic assessment moves on to genetic testing. In families who need genetic testing, it is important that the best person is tested. Certain signs, like the age people were when they first had cancer, help to decide the best person in a family to test. The person who has the assessment may not be the person who is tested. Your health care provider can contact the Maritime Medical Genetics Service with any questions.

**Increased risk for ovarian cancer:** The risk of ovarian cancer after breast cancer is highest in women with a family history of breast cancer. A strong family history of breast cancer may be caused by an inherited genetic change (usually in the BRCA1 or BRCA2 gene). These genetic changes can also cause ovarian cancer. If there is no genetic cause for breast cancer then the risk of ovarian cancer is not elevated.

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**Notes and Questions**
Returning Cancer: Signs to Watch For

One of the biggest worries for cancer patients is that the cancer may come back. If the cancer does come back, it is usually in the first three to five years. This is why the follow up care is so important. After five years, the chance of cancer coming back drops every year.

Your follow up care will focus on finding cancer that comes back in the breast or lymph nodes. It is possible to treat the cancer if it is found early.

Bone scans and CT scans are not usually part of your follow up care. These scans may be done if you have any signs that suggest cancer has come back. See the list below for some of these signs.

Please notify your health care provider if you have any of the following:

- A new pain that won’t go away.
- A cough that won’t go away.
- A lump in your breasts, neck, armpit or any other part of your body.
- Unusual changes in the site of your surgery or in the scar itself.
- A tired feeling that won’t go away.
- Loss of appetite.
- Weakness, tingling or numbness in the arm, hand or leg.
- Swelling in the arm, shoulder, breast, chest, armpit or back.
- Any new symptom that is unusual or severe and doesn’t go away, such as bone pain, shortness of breath or headaches.

Notes and Questions
Cancer Screening and Other Tests

Just like people who have not had breast cancer, it is important for you to continue with routine screening for cancer and other diseases. Some examples are given below but others may be needed depending on your family history. Talk with your health care provider about cancer screening and other tests.

Cervical cancer screening: Women should start screening for cervical cancer (also called a Pap test) within three years of becoming sexually active or upon reaching the age of 21, whichever comes later. Women who have never been sexually active do not need Pap screening. Pap testing should be done every three years. Women with a history of cancer of the cervix or who have been treated (by LEEP, laser, cryotherapy, cone, hysterectomy) should be screened annually for life. Immunocompromised or HIV positive women should be screened annually for life. Women can discontinue having Pap tests at age 70 if they have had three or more negative Pap tests in the previous 10 years. Women who have had a total hysterectomy for benign causes with no treatment for abnormal cell growth on the cervix can discontinue Pap screening.

The screening guidelines only apply to women with no symptoms of cervical cancer who are, or have been, sexually active. If a woman is experiencing symptoms such as irregular bleeding or pain, she should speak with her health care provider. To learn more about cervical cancer screening guidelines, call Nova Scotia Cancer Care Program: 1-888-480-8588 or visit www.nscancercare.ca.

For Men

Prostate Cancer: Talk to your health care provider about your risk for prostate cancer and the benefits and harms of PSA testing. To learn more visit the Canadian Cancer Society’s website www.cancer.ca or call 1-888-939-3333.

For Both Men and Women

The Nova Scotia Colon Cancer Prevention Program: All Nova Scotian women and men, who are between 50 and 74 years old and are registered with MSI (have a Nova Scotia Health Card), will automatically get a colon cancer home screening test in the mail every two years. Colon cancer screening works best when you take the test when you are feeling good – before you have any symptoms or warning signs. Talk with your health care provider if you have concerns about colon cancer or are having long-lasting changes in your bowel habits. To learn more about the colon cancer home screening test call Nova Scotia Cancer Care Program: 1-866-599-2267 or visit www.nscancercare.ca.

Continued on next page
Cancer Screening and Other Tests (continued)

For Both Men and Women (continued)

Diabetes screening: Starting at age 40, you should have a Fasting Plasma Glucose (FPG) test every 3 years. For those at high risk (family history, overweight, Aboriginal, African Nova Scotian, have a history of gestational diabetes or pre-diabetes, heart disease, increased cholesterol, increased blood pressure), more frequent and/or earlier testing should be considered. To learn more, call the Canadian Diabetes Association 1-800-226-8464 or visit [www.diabetes.ca](http://www.diabetes.ca).

Cholesterol screening: You should have a cholesterol test at age 40 if you are a man, or age 50 if you are a woman. Start at a younger age if you have diabetes, heart disease, if you smoke or if you are overweight. Talk with your health care provider about how often you should have this test.

Blood pressure checks: You should have your blood pressure checked regularly by your health care provider.

Notes and Questions
Other Ways to Protect Your Health

**Immunizations:** You should have a flu shot every fall. If you are 65 or older, or have diabetes, heart or lung problems, you should also have a pneumonia vaccine. Ask your health care provider about the benefits and risks of getting a shingles vaccination.

**Protect your skin:** Having chemotherapy or radiation can make you more sensitive to harm from the sun and other sources of ultraviolet (UV) rays. Some prescription and over-the-counter medications can also make skin very sensitive to the sun. Protect yourself by being informed and using good sun protection practices at work, in your community and on vacation. Check the UV Index daily and practice sun safety as recommended. The higher the UV Index, the more protection is required. Plan activities in the shade, cover up with clothing and a wide-brimmed hat, wear sunglasses and use a broad spectrum sunscreen with a Sun Protection Factor (SPF) of 30 or higher. Avoid use of tanning beds.

**Dental health:** Chemotherapy can cause dental problems. It is important that you have regular dental check-ups. Your dentist will need to know that you had breast cancer and the type of treatments you received. Brushing with a soft tooth brush, and using toothpaste for sensitive teeth is recommended.

**Physical activity:** Whether you are getting back to activities you enjoy or starting new ones, being active after treatment for cancer is important. Physical activity can help improve appetite, energy level and sleep. Physical activity can also help reduce risk of cancer coming back, fatigue and anxiety. It is recommended you try to be physically active at least 150 minutes per week and that you should start slowly and progress gradually. It is important to always check with your health care provider about the activities and level of activity that would be best for you. For more information on how to get active visit, [www.phac-aspc.gc.ca](http://www.phac-aspc.gc.ca) and [www.heartandstroke.com](http://www.heartandstroke.com).

**Healthy eating:** There are no special foods or vitamins to take after cancer treatment. Following Canada’s Food Guide is the best approach [www.hc-sc.gc.ca/fn-an/index-eng.php](http://www.hc-sc.gc.ca/fn-an/index-eng.php). A clinical dietician can help you learn more about eating well.

For the QEII Cancer Care Program please call the clinical dietician at **902-473-3972**.

For the Cape Breton Cancer Centre please call the clinical dietician at **902-567-8552**.

Call your cancer patient navigator **(1-866-524-1234)** for referral to a dietician in your community.
What to Expect after Cancer Treatment

Most people have some side effects after breast surgery and/or breast cancer treatment. Side effects can start right away or they can take weeks to months to begin. Some side effects improve with time, while others may go on for many years or may be permanent. These side effects may be physical or emotional.

If any of these side effects are a problem for you, talk to your health care provider. If there is a cancer patient navigator in your area, they can also help. Call 1-866-524-1234.

Your health care provider or cancer patient navigator should be able to provide you with support, connect you with other services and give you information about local resources.

If you visit a clinic with a group practice, it is best if one health care provider is responsible for your follow up care.

Cancer-related fatigue: Fatigue (excessive tiredness) is very common after breast cancer treatment and can last a long time. Your health care provider will assess your fatigue and suggest ways to prevent or manage it. Regular physical activity is one of the best ways to decrease your fatigue. There are resources available to help you learn more about cancer-related fatigue. For more helpful hints visit the websites listed at the end of this booklet.

Brain fog or chemo brain: Some people notice they are forgetful, have trouble focusing, have memory problems or are confused after being diagnosed and treated for breast cancer. Exercising your brain by doing crossword puzzles or number games will help. For more helpful hints visit the websites listed at the end of this booklet.

Depression: Depression may occur after cancer treatment. Continuing to be active with your family, and in your community, can help you deal with depression. Physical activity and hobbies are also helpful. Your health care provider may suggest other approaches, such as medication or talking to a counselor. The Depression Hurts website has information you may find helpful, www.depressionhurts.ca. Your family members may also have issues with depression, anxiety and mood swings. They, too, will benefit from contacting their health care provider.

Difficulty Returning to work: Many people experience distress at the idea of returning to work after treatment. When you do go back, you may find your work more challenging than before cancer treatment. Talk with your employer, your workplace’s employee health staff and your health care provider about how to deal with your work related concerns.

Continued on next page
What to Expect after Cancer Treatment (continued)

**Lymphedema:** Arm or neck swelling (lymphedema) can occur after surgery or after radiation treatment. Signs of lymphedema include swelling or tightness of your arm, neck, shoulder or chest; a feeling of heaviness or aching in your arm. A compression garment may be used to decrease the swelling.

Moderate physical exercise of your arm and moderate weight lifting or resistance training do not increase your risk of developing lymphedema. In fact, all these activities can be helpful.

If you develop lymphedema, you can be referred to the Lymphedema Clinic in Halifax by your health care provider or you can self-refer by calling the Clinic at 902-473-2151.

If you prefer to be seen in Sydney, a referral needs to be made to the Physiotherapy Department at the Cape Breton Cancer Centre. Referrals can be from physicians or any other health care provider. For more information call 902-567-7748 and the fax number is 902-567-7869.

**Shoulder stiffness:** This is usually related to surgery to remove your lymph nodes in your armpit or radiation in that area. Gentle exercise may help. Your health care provider or local pharmacist can suggest other options if the stiffness continues.

**Aches and pains:** You may have general body aches as a side effect of your treatments. This may last for a time after your therapy is completed. Usually acetaminophen, heat or warm baths and gentle exercise will help. If your aches and pains continue to interfere with your ability to enjoy yourself, your health care provider can suggest other options.

**Trouble sleeping, anxiety, mood swings:** It is not uncommon to have emotional distress after cancer treatment. Your health care provider will be able to give you suggestions to manage your emotional distress.

**Early or treatment related menopause:** Women should talk to their health care provider about any changes in their periods. Cancer treatment may cause women to go into menopause or start at a younger age than usual. Women may have hot flashes, vaginal dryness, mood changes or changes in how they respond sexually. Health care providers can help with these symptoms or suggest someone to talk to.

**Sexual problems:** Cancer treatment may cause women to go into menopause which can cause sexual problems. You may have pain during sex, vaginal dryness, decreased or loss of sexual interest, decreased ability to become sexually aroused or relationship difficulties. Your health care provider may be able to give you guidance and support or suggest another health care provider who can.

**Weight gain:** Weight gain is not unusual after cancer treatment. Having healthy weight, eating well and exercise are all important for your overall health. (See page 9 for more information).

**Body Image Changes:** After treatment and surgery for breast cancer, it is not uncommon for people to experience changes in their body image. Useful advice can be found in the Support & Resources section beginning on page 13. Discussions with your health care provider can also be helpful.
Programs for Cancer Survivors

Cancer Transitions

The Cape Breton Cancer Centre offers a free six week program, Cancer Transitions, to help cancer survivors adjust to life after cancer.

Cancer Transitions covers these topics:

- Get Back to Wellness
- Exercise for Wellness
- Eating Well
- Medical Care After Cancer
- Emotional Health
- Exercise for Wellness

For more information and to register:
Cape Breton Cancer Centre call 902-567-8074

Living Beyond Cancer: What Happens Next?

The Nova Scotia Health Authority offers a two and a half hour session that focuses on some of the key issues cancer survivors face in the months following active treatment. You may attend in person at the Bethune Ballroom, VG Site and at the Cape Breton Cancer Centre or via telehealth, at the Regional Hospital in your area.

Topics include:

- What is Follow up Care
- Eating Well After Cancer
- What to Expect after Cancer Treatment
- Physical Activity and Emotional and Practical Support.

For more information and to register call toll free 1-866-599-2267 and select option 3, or to register online please go to www.nscancercare.ca and click on I am a Cancer Survivor.

We now have Living Beyond Cancer video segments available to watch online. These can be found by visiting library.nshealth.ca/cancer and clicking on Cancer Patient Education Videos.

Your Way to Wellness

Your Way to Wellness is a provincially sponsored, free program, led by volunteer lay leaders. The Program is designed to help people with chronic conditions learn skills to live a healthier life. Your Way to Wellness is offered once a week for a 2 ½ hour session over six weeks. Programs are offered all over Nova Scotia.

For more information, call toll free, 1-888-672-3444 or visit www.yourway.novascotia.ca

Continued on next page
Programs for Cancer Survivors (continued)

Group Learning Sessions for Cancer Patients and Families

Some cancer patients and survivors experience treatment related side effects. There are a number of group teaching sessions which focus on the most common ones.

These sessions are not support groups, but participants have the opportunity to ask questions. Each session is 2 hours. Available resources and supports are discussed. Family and friends are also welcome to attend. For the date, time and location of upcoming sessions, please call 1-866-599-2267 and select option 3.

We now have video segments available to watch online. These can be found by visiting library.nshealth.ca/cancer and clicking on Cancer Patient Education Videos.

Managing Your Cancer-related Fatigue: Cancer-related fatigue (tiredness) is one of the most common side effects of cancer treatment. The sessions provide practical information about cancer-related fatigue and helpful management strategies.

Taking Control of Your Cancer-related Worry and Anxiety: Most people with cancer experience worry and some level of anxiety. This session provides practical information about cancer-related anxiety and helpful coping strategies (such as deep breathing relaxation).

Taking Control of Your Cancer-related Sadness and Depression: Many people with cancer experience sadness or depression. This session provides practical information about cancer-related sadness and depression and helpful coping strategies.

Coping With Cancer-related Brain Fog: Some cancer patients and survivors notice changes in their thinking and cognitive abilities during or after treatment. This is known as cancer-related brain fog. This session focuses on understanding cancer-related brain fog and management strategies.

*Coping with Your Cancer-related Pain: Not everyone with cancer experiences pain but many people do. This session focuses on understanding cancer-related pain and management strategies such as medication, managing energy and relaxation.

*This session is only available online at library.nshealth.ca/cancer and click on Cancer Patient Education Videos.
Support & Resources

Cancer affects more than your body. It can have an impact on your whole life. Many people who have had cancer find that talking with a professional counselor or therapist can be very helpful. Health care providers are the experts in treating cancer, but you are the expert in the way cancer affects your life. Counselling can help you to find strength and meaning. It can also help you cope, adjust, and find solutions to problems. You can go to counselling on your own or with members of your family.

The QEII Cancer Care Program has a psychosocial cancer team for people who have been diagnosed and treated for breast cancer anywhere in the province of Nova Scotia. The team provides counselling to breast cancer patients & their families. Team members include psychologists, psychiatrists, social workers, nurses and spiritual care providers.

Support can be provided to you by phone or in person. If you are having difficulty coping, you can ask your health care provider, your cancer doctor or another member of your cancer team to make a referral to a member of the psychosocial cancer team (QEII Cancer Care Program).

To reach the social worker at the Cape Breton Cancer Centre please call 902-567-8551.

If you are unable to travel to Halifax or Sydney, check with your health care provider to find out about public and private services in your district. To reach your local cancer patient navigator please call 1-866-524-1234.

Cancer Connection – Canadian Cancer Society

The Canadian Cancer Society can connect patients or caregivers with fully screened and trained volunteers who’ve had cancer or cared for someone with cancer. Volunteers listen, provide hope, offer encouragement, and share ideas for coping – all from their unique perspective as “someone who has been there.” Call 1-888-939-3333 to get connected or logon at www.cancerconnection.ca.

Private Medical Insurance and Employee Assistance Programs

Counselling is also covered under some private insurance plans (Blue Cross). There may be a yearly maximum as to how much your plan will pay. Contact your insurer for the details specific to your plan.

As well, many employers offer Employee Assistance Plans which offer a limited number of free counselling sessions. Please contact your Human Resources Department for information.

Continued on next page
Support & Resources (continued)

Support Groups

A support group is a group of people with similar problems or concerns. Support groups meet on a regular basis. There are many different cancer support groups. Some are for the patients. Others are for the family and/or friends of the patient. Others are open to anyone. Groups may be led by a health care professional or made up completely of peers. You can find support groups that meet face-to-face or online. It may help to talk with the person running the group to see if the group offers what you need. You may want to check out a few different groups to find one that works best for you. It is hard to know if a support group will be right for you without going to at least two meetings. A support group should make you feel comfortable enough to talk about what you think and feel. If it does not, speak with the facilitator privately. You may decide it is best not to continue going. Remember that it may take time to find a group that fits.

Find a support group in your area:

- Canadian Cancer Society: 1-800-639-0222 or visit www.cancer.ca
- QEII Cancer Care Program in Halifax: 902-473-6067
- Social Worker at the Cape Breton Cancer Centre, Sydney: 902-567-8551
- Cancer Patient Navigators: 1-866-524-1234
- Check for posters in your cancer treatment centre.
- Ask people you know who have gone through, or who are currently dealing with cancer themselves.

Many people choose online support groups. These groups help people who may not wish to attend a face-to-face group. Be careful when looking for an online group. Anyone with internet access can create an online group, even if they do not have any qualifications.

If you are interested in online groups, contact Cancer Chat Canada, by visiting their website, cancerchatcanada.ca. Canadian cancer care professionals lead these groups. Most are available in any province or territory.

You can also call 211 to find local support groups.

Books

After You Ring the Bell...10 Challenges for Cancer Survivor. Anne Katz. Oncology Nursing Society. 2012.

Cancer is a Word Not a Sentence by Dr. Robert Buckman.
Support & Resources (continued)

Books (continued)


The Essential Cancer Treatment Nutrition Guide and Cookbook by Jean Lamantia

The Healing Circle by Dr. Timothy Walker and Dr. Rob Rutledge. 2011.

The Healing Journey: Overcoming the Crisis of Cancer by Alastair J Cunningham. 2010.

The Places that Scare You: A Guide to Fearlessness in Difficult Times by Pema Chodron


When a Parent is Sick: Helping Parents Explain Serious Illness to Children. Joan Hamilton, Clinical Nurse Specialist, Cancer Care, QEII, Halifax.


Websites

The Web is a great source of information about moving forward after cancer, but it is also full of misinformation. You can rely on these websites:

For Breast Cancer Survivorship information, visit:
- American Society of Clinical Oncology (ASCO) Patient Site: www.cancer.net
- British Columbia Cancer Agency: bccancer.bc.ca
- Canadian Breast Cancer Foundation: www.cbcf.org or call 1-866-273-2223

Continued on next page
Support & Resources (continued)

Websites for Breast Cancer Survivorship information (continued)

- Canadian Breast Cancer Network: www.cbcn.ca or call 1-800-685-8820
- Canadian Cancer Society: www.cancer.ca <click on> Nova Scotia or call 1-888-939-3333
- Cancer.net (breast cancer male): www.cancer.net
- Healing and Cancer: www.healingandcancer.org
- Living Beyond Cancer: www.nscancercare.ca <click on> Patients and Families <click on> I am a Cancer Survivor
- National Comprehensive Cancer Network: www.nccn.org/patients/resources/life_after_cancer
- National LGBT Cancer Network: www.cancer-network.org
- Rainbow Health Ontario: www.rainbowhealthontario.ca

- Rethink Breast Cancer: www.rethinkbreastcancer.com
- Willow: www.willow.org

For Physical Activity information, visit:

For Fatigue information, visit:
- Canadian Cancer Society: www.cancer.ca <click on> Cancer Information <click on> diagnosis and treatment <click on> managing side effects <click on> fatigue
- American Cancer Society: www.cancer.org/treatment/treatments-and-side-effects/physical-side-effects/fatigue
- www.youtube.com/watch?v=YTFPMYGe86s&feature=youtu.be

For Sexual Health information, visit:
- Canadian Cancer Society: www.cancer.ca <click on> Cancer Information <click on> diagnosis and treatment <click on> managing side effects

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Support & Resources (continued)

For Information in Languages other than English
There are a number of reputable sources of cancer information in languages other than English. We suggest the following Canadian resources:
- British Columbia Cancer Agency: bccancer.bc.ca
- Canadian Cancer Society: www.cancer.ca or call 1-888-939-3333
- Cancer Care Manitoba: www.cancercare.mb.ca
- Cancer Council New South Wales: www.cancercouncil.com.au
- Vancouver Coastal Health: http://vch.eduhealth.ca

Get Involved

Cancer patients, families, and friends have unique, valuable perspectives to share about cancer care. It is important to hear from people who have experience with the cancer system in order to see what can be changed to better meet people’s needs.

As a cancer survivor you may want to get involved with the Cancer Patient Family Network (CPFN). The CPFN is a program of Nova Scotia Cancer Care Program that connects cancer patients, survivors, family members, and friends with opportunities to improve the cancer system. As a member of the CPFN you will have opportunities to participate in focus groups and committees, complete surveys, and review patient education resources.

Contact Nova Scotia Cancer Care Program by calling toll-free 1-866-599-2267, emailing cancercareinfo@nshealth.ca or visiting www.nscancercare.ca. For more information about the Cancer Patient Family Network, please contact the Patient Engagement Coordinator at 902-473-2637 or email CPFN@nshealth.ca.
Follow up Care Schedule
You can use these charts to keep track of your follow up care.

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<th>Family Doctor/Nurse Practitioner’s visit</th>
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Notes and Questions

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References


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