Follow up Care for Colon Cancer Patients

Living Beyond Cancer
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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.

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.

Inherited risk of colorectal cancer: Only 5-10% of colorectal cancer is hereditary (passed down through a family). However, it is important to identify individuals who could be at risk of inherited colorectal cancer as their screening and management options would be different from a person who developed colorectal 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 often based on family history information and your own personal medical history it is important to have accurate information about the cancer in your family. 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. The best place to start is with a discussion with your health care provider. Your health care provider can contact the Maritime Medical Genetics Service with any questions.

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 your cancer treatment or fear of recurrence.

- **Check for return of your cancer or a new cancer.** Early detection of cancer is important. Treatment is more likely to be successful at an early stage.

Follow up care for colon 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. Your surgeon may also be involved in providing your follow up care.

- A **blood test called CEA** that may help find a return of cancer earlier.

- A **Colonoscopy** test that allows a doctor to look for polyps or second cancers with a scope (lighted tube).

- **X-rays or scans** For example: X-ray, CT scan or ultrasound. X-rays may be suggested for the chest, abdomen and the pelvis.

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 Recommendations

For Colon Cancer by Years after End of Treatment

If you are diagnosed with a genetic syndrome linked with colon cancer you may have different follow up. Please discuss with your surgeon or medical oncologist.

<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>
</thead>
<tbody>
<tr>
<td>Health Care Provider Visit</td>
<td>Every 3 to 6 months</td>
<td>Every 3 to 6 months</td>
<td>Every 3 to 6 months</td>
<td>Every 6 months</td>
</tr>
<tr>
<td>CEA test</td>
<td>Every 3 to 6 months</td>
<td>Every 3 to 6 months</td>
<td>Every 3 to 6 months</td>
<td>Every 6 months</td>
</tr>
<tr>
<td>CT Scan</td>
<td>A CT of your chest, abdomen and pelvis should be done annually (depending on your overall health).</td>
<td>When your health care provider feels it is needed</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

After 5 years, you and your health care provider will discuss the need for future visits and tests.

Colonoscopy Follow up by Years after Diagnosis

To be coordinated with the attending surgeon.

<table>
<thead>
<tr>
<th>Colonoscopy</th>
<th>Year 1</th>
<th>Year 3</th>
<th>Year 5</th>
<th>Ongoing</th>
</tr>
</thead>
<tbody>
<tr>
<td>A complete colonoscopy should be done around the time of diagnosis.</td>
<td>A complete colonoscopy should be done one year after diagnosis.</td>
<td>If previous colonoscopy shows no signs of a recurrent tumour or polyps, a colonoscopy should be done at three years.</td>
<td>If previous colonoscopy shows no signs of a recurrent tumour or polyps, a colonoscopy should be done at five years.</td>
<td>If normal every 3 to 5 years thereafter, as long as you are in good health</td>
</tr>
</tbody>
</table>

Notes and Questions

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Returning Cancer: Signs to Watch For

One of the biggest concerns for cancer patients is that the cancer may return some day. If colon cancer does return, it is usually in the first three to five years after diagnosis. This is why the follow up care is so important. After five years, the chance of cancer returning drops year after year.

If your cancer returns, it is usually because cancer cells have spread to the liver or lungs. There is also a small risk of the cancer coming back in the bowel where the surgery was done or a new cancer growing in a different part of the bowel. This is why the follow up care has tests for the liver, lungs and colon. It is possible to treat the cancer if it is found early and can be removed. If it cannot be removed, cure is usually not possible, but chemotherapy will improve symptoms and may extend your life.

Know the signs to watch for. Please notify your health care provider if you have any of these changes in your health:

Notify your health care provider right away if you see:
- Blood in your bowel movement
- Blood that you cough up
- Bowel movement that is very pale or black
- Yellow eyes or skin (jaundice)
- New lumps, especially in the neck and groin
- Blood in your urine
- If you have an ostomy, a change in your stoma

See your health care provider if these symptoms last more than one week:
- A change in your bowel movements, like loose bowel movements or constipation
- Cough
- Pain in your abdomen or rectal area (or where the rectum/anus used to be)
- Nausea (feeling sick to your stomach)
- Pain that will not go away or is getting worse
- Vomiting that lasts more than a few days
- Shortness of breath
- Loss of appetite
- Weight loss without trying
- Drop in your energy level or ability to be active
Living Beyond Cancer

Follow up Care for Colon Cancer Patients

Cancer Screening and Other Tests

Just like people who have not had colon 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.

For Women

Cervical Cancer Screening: Pap tests should begin 3 years after becoming sexually active or upon reaching the age of 21 – whichever comes later. If you have never been sexually active you do not need to have Pap tests. You should have your Pap test every three years. You can discontinue having Pap tests at age 70, only if you have a history of negative Pap test results. If you have had your cervix removed through a hysterectomy (not for cancer), you do not need to have Pap tests. Immunocompromised or HIV positive women and women who have been treated for cervical dysplasia or have a history of cancer of the cervix should have annual screening for life. Talk with a doctor or health care provider if you have any concerns about cervical cancer or are experiencing symptoms. To learn more about cervical cancer screening guidelines, call Cancer Care Nova Scotia: 1-888-480-8588 or visit www.cancercare.ns.ca.

Breast Cancer Screening: Women 50 and over should have a mammogram every 2 years. If you have a strong family history or are currently using hormone replacement therapy a mammogram may be needed every year. Visit the Nova Scotia Breast Screening Program website www.breastscreening.nshealth.ca to learn more. Call 473-3960 or 1-800-565-0548 to book an appointment.

For Men

Prostate Cancer: Talk to your doctor/nurse practitioner 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

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.

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

For Both Men and Women (continued)

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: Get your blood pressure checked regularly by your health care provider.

Other ways to protect your health:

Immunization: You should have a flu shot every fall. If you are 65 or older, 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.

Sun Safety: It is important to use sun protection. Use sunscreen (minimum SPF 30), wear a wide brimmed hat, long sleeved tops, long pants and sunglasses.

Stop Smoking: Smoking may affect your recovery from treatment, increase the risk of your cancer returning or getting another cancer. Helpful resources can be found at http://www.gov.ns.ca/hpp/addictions/tobacco/makesyousick.asp and www.smokershelpline.ca

Limit Alcohol Use: Excessive alcohol use may increase the risk of your cancer coming back or getting a new cancer. A helpful resource can be found at http://www.gov.ns.ca/hpp/addictions/alcohol/.

Dental Health: Chemotherapy can cause dental problems. It is important that you have regular dental checkups. Your dentist will need to know that you had colon cancer and the type of treatments you have received. Brushing with a soft tooth brush, and using toothpaste for sensitive teeth is recommended.
What to Expect after Cancer Treatment

You may have some side effects after your surgery and/or cancer treatment. Some side effects can start right away or can take 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 can 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 only one health care provider is responsible for your follow up care.

<table>
<thead>
<tr>
<th>Bowel Problems</th>
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<tbody>
<tr>
<td>Urgent need to have a bowel movement</td>
</tr>
<tr>
<td>Loss of bowel control (incontinence)</td>
</tr>
<tr>
<td>Diarrhea</td>
</tr>
<tr>
<td>Constipation</td>
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<tr>
<td>Frequent bowel movements</td>
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**What to do:** Talk to your health care provider.

<table>
<thead>
<tr>
<th>Bowel Blockage</th>
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<td>(pain, nausea, vomiting, not able to pass gas or bowel movements)</td>
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**What to do:** This is an emergency, go to the emergency department right away.

<table>
<thead>
<tr>
<th>Ostomy Issues</th>
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<tbody>
<tr>
<td>Skin or appliance problems</td>
</tr>
<tr>
<td>Constipation, diarrhea</td>
</tr>
<tr>
<td>Irrigation problems</td>
</tr>
<tr>
<td>Hernia – A hernia is a weakness in the wall of the abdomen that allows the contents to bulge out forming a lump or swelling. When you have a stoma, a hernia can change how the Ostomy appliance fits and you may need to change products or use a hernia support binder.</td>
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**What to do:** Talk to your Entrostomal Therapy Nurse, your health care provider or cancer patient navigator.

<table>
<thead>
<tr>
<th>Urination Problems</th>
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</thead>
<tbody>
<tr>
<td>Frequent urination</td>
</tr>
<tr>
<td>Difficulty emptying your bladder</td>
</tr>
<tr>
<td>Urgent need to urinate</td>
</tr>
<tr>
<td>Incontinence (loss of bladder control)</td>
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</table>

**What to do:** Contact your health care provider or cancer patient navigator.

<table>
<thead>
<tr>
<th>Pain in the Pelvis</th>
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<td><strong>What to do:</strong> Contact your health care provider or cancer patient navigator.</td>
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<tr>
<th>Nerve Damage/Neuropathy</th>
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<tr>
<td>Tingling, numbness or pain in the hands or feet</td>
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**What to do:** Contact your health care provider or cancer patient navigator.

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

Sexual Problems
Men
- Problems with sexual function (getting or keeping an erection, changes in ejaculation)
- Decreased or loss of sexual interest/desire
- Relationship difficulties

Women
- Pain during sex
- Decreased or loss of sexual interest/desire, decreased ability to become aroused
- Vaginal dryness
- Relationship difficulties

What to do: Contact your nurse where you received your cancer care or the cancer patient navigator. They can help you find resources such as the psychosocial oncology team and couples counselling. See the resources at end of this guide.

Fatigue and Less Energy
- Extreme tiredness
- Weakness
- Usual activity feels harder to do
- Difficult to do physical activity

What to do: Tell your health care provider who will assess your fatigue and suggest ways to prevent or manage it. Regular exercise is one of the best ways to decrease your fatigue. There are resources available to help you learn more about your cancer related fatigue. For more helpful hints visit Canadian Partnership Against Cancer: www.cancerview.ca <click on> idc <click on> groups <click on> public <click on> documents <click on> webcontent <click on> manage_cancer_fatigue.pdf.

Difficulty Sleeping
- Difficulty falling asleep
- Waking up through the night and not being able to go back to sleep

Return 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 your cancer treatment.

What to do: Talk with your employer, your workplace’s employee health staff and your health care provider about how to deal with your work related concerns.

Psychosocial Distress
- Trouble sleeping
- Memory and concentration problems
- Mood swings
- Emotional distress
- Fear of cancer coming back
- Anxiety
- Depression

What to do: Contact your nurse where you received your cancer care or the cancer patient navigator in your area (if available). They can help you find resources such as the psychosocial oncology team and social workers in your area. See the resources at end of this guide.
Eating Well after Colon Cancer

Colon cancer survivors may have problems when they start to eat fruits, vegetables, whole grains, legumes and other high fibre foods again. Each cancer survivor is unique. It is important to discuss your issues with your doctor or a dietician.

For the Nova Scotia Cancer Centre please call the clinical dietitian at 902-473-3972. For the Cape Breton Cancer Centre please call the clinical dietitian at 902-567-8552. Call your community Cancer Patient Navigator 1-866-524-1234 for referral to a dietitian in your community.

For patients with ostomies who have questions or concerns about their ostomy, please contact the Enterostomal Therapy Department at:
Capital Health 902-473-7503
Cape Breton District Health Authority call 902-567-8000 ext. 2335
Check with your community cancer patient navigator (1-866-524-1234) for referral to an Entrostomal Therapy Nurse in your district if you do not already have one.

The tips below may help you with side effects of your surgery and/or treatment. Please discuss with your surgeon, Enterostomal therapist or dietitian to determine what is best for you.

- Eat smaller, more frequent meals.
- When trying new foods, start with very small servings.
- Drink at least 6 - 8 cups (1.5 - 2 litres) of fluid per day.
- Drink fluid between meals rather than with meals.
- Limit drinks with caffeine (such as coffee or cola soft drinks).
- Limit alcohol.
- Limit fatty or greasy foods (such as french fries, fried meats, bacon, potato chips, gravies and rich desserts).
- Limit spicy foods.
- Gradually eat more foods that are high in soluble fibre such as oatmeal, barley, white rice, bananas, white bread, applesauce and canned fruit such as peaches and pears.
- Eat less food with insoluble fibre such as wheat bran, whole grain breads and cereals, beans, peas, popcorn, any bran products and raw vegetables.
- Eat well cooked vegetables.
- Limit how much red meat you eat.
- Peel and remove seeds of fruits and vegetables.
- To reduce gas, limit these foods: cabbage, onions, dried beans and peas, lentils, lettuce, cucumber, broccoli, cauliflower, radishes, brussel sprouts, corn, turnip, green pepper, sauerkraut, melons, grapes, raw apple, prunes, raisins.
- To reduce gas, avoid chewing gum, talking while eating, drinking carbonated beverages (pop) and drinking with a straw.

Once you feel better, slowly start to put foods back into your diet in small amounts, one at a time. If you get diarrhea, gas or cramping after re-starting a food, avoid that food and try it again at a later time.
Physical Activity after Colon Cancer

Whether you are getting back to activities you enjoy or starting new ones, being active after treatment for colon cancer is important. Research has suggested a link between physical activity and decreased rates of cancer return and increased rates of survival in colon cancer. Physical activity can also help with fatigue and sleep disturbances. Always check with your doctor/nurse practitioner before starting an exercise program.

Following treatment, you may need to take special precautions if you:
- have anemia, a ‘low red blood cell count’.
- have problems with your balance.
- have numbness or tingling in your hands or feet.
- are taking any pain or anti-inflammatory drugs. These drugs may mask pain so be careful to not “over-do-it” and take care of any injuries immediately.
- have weakened immune function.
- have had surgery in your abdominal cavity since heavy lifting can cause a hernia.

Be sure your incision has completely healed before gradually introducing weights.
While lifting weights be sure to concentrate on breathing with exertion and do not hold your breath or bear down.

Physical activity can speed up your digestion. This can increase the need to go to the bathroom during or immediately after activity. Follow these helpful hints:
- Go to the bathroom before your activity and take short breaks to go to the bathroom.
- Know where the bathrooms are located along the way.
- An urgent time is usually right after you stop, so have easy access to the bathroom at that time.
- Wear clothing that can be quickly and easily removed.
- If you are using a medication for diarrhea, take it prior to exercising.
- Sip water during your activity – do not gulp down large amounts.
- The harder the activity or the more you are sweating, the more water you should drink.
- If you are worried about having an ‘accident’ take an extra set of clothes along with you, as well as a wash cloth and a plastic bag for soiled clothes.

Doing physical activity when you have an Ostomy can be hard, at first. Talk with your Enterostomal Therapy Nurse about your activity and they will help you. There may be specific supplies and techniques that can assist with the issues you are having, such as protection and securing of appliances.
Programs for Cancer Survivors

**Cancer Transitions**

Capital Health Cancer Care Program and the Cape Breton Cancer Centre offer 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
- Emotional Health
- Eating Well
- Medical Care After Cancer

For more information and to register:
Capital Health Cancer Care Program call 902-473-8241
Cape Breton Cancer Centre call 902-567-8074

**Living Beyond Cancer: What Happens Next?**

The Capital Health Cancer Care Program 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 or via telehealth, at the Regional Hospital in your area.

**Topics include:**
- What is Follow up Care
- Eating Well After Cancer
- Physical Activity and Emotional and Practical Support.

For more information and to register, call toll free: 1-866-524-1234 or to register online please go to [www.cancercare.ns.ca](http://www.cancercare.ns.ca)

We now have Living Beyond Cancer video segments available to watch online via our YouTube channel. These can be found by visiting [www.cancercare.ns.ca](http://www.cancercare.ns.ca) and clicking on Patients and Families.

**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.yourway2wellness.gov.ns.ca](http://www.yourway2wellness.gov.ns.ca)
Support and Resources

Cancer affects more than your body. It has 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.

There is a psychosocial cancer team at the Nova Scotia Cancer Centre for people who have been diagnosed and treated for colon cancer anywhere in the province of Nova Scotia. The team provides counselling to colon 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. Although you may be medically discharged from the Cancer Program, the psychosocial cancer team is still available to help you. For more information, call: 902-473-6067 (Nova Scotia Cancer Centre).

If you live in Sydney, please call the social worker at the Cape Breton Cancer Centre at 902-567-8551. If you live outside Halifax and Sydney, check with your cancer patient navigator, social worker or health care provider to find out about 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 have had cancer or cared for someone with cancer. Our 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.

Colorectal Cancer Association of Canada – Cancer Coaches

Cancer Coaches are volunteers with the Colorectal Cancer Association of Canada who share their experiences and information in various aspects of colorectal cancer. They can help address your concerns and answer your questions.

Login at http://www.colorectal-cancer.ca/en/find-support/cancer-coaches/

Private Medical Insurance and Employee Assistance Programs

Counselling is also covered under some private insurance plans (i.e. 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.

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.

Continued on next page
Support and Resources (continued)

Support Groups (continued)
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 at 1-800-639-0222 or visit www.cancer.ca.
- Ostomy Canada Society has 4 chapters in NS that run support groups. Visit www.ostomycanada.ca or call 1-888-969-9698.
- Social Worker at the Cape Breton Cancer Centre Sydney: 902-567-8551.
- 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 real experience.

If you are interested in online groups, contact CancerChatCanada by visiting their website at 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
The Essential Cancer Treatment Nutrition Guide and Cookbook by Jean Lamantia
The Healing Circle by Dr. Timothy Walker and Dr. Rob Rutledge, 2011.
The Places that Scare You: A Guide to Fearlessness in Difficult Times by Pema Chodron

Continued on next page
Support and Resources (continued)

Websites

The Web is a great source of information about moving forward after cancer, but it is also full of misinformation. These websites are ones you can rely on!

For Colon Cancer Survivorship information, visit:
- American Society of Clinical Oncology (ASCO) Patient Site: www.cancer.net
- British Columbia Cancer Agency: bccancer.bc.ca
- Canadian Cancer Society: www.cancer.ca <click on> Nova Scotia or call 1-888-939-3333
- Cancer Care Nova Scotia: www.cancercare.ns.ca
- Colorectal Cancer Association of Canada: www.colorectal-cancer.ca or call 1-877-50-COLON
- Healing and Cancer: www.healingandcancer.org
- Ostomy Canada Society: www.ostomycanada.ca or call 1-888-969-9698

For Physical Activity information, visit:
- www.participation.com
- www.heartandstroke.com/site/c.iKIQLcMWJtE/b.3484253/k.4FE0/Healthy_living__Physical_Activity.htm

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.
- Canadian Partnership Against Cancer: www.cancerview.ca/ <click on> idc/<click on> groups/<click on> public/<click on> documents/<click on> webcontent/<click on> manage_cancer_fatigue.pdf.
- American Cancer Society: www.cancer.org <click on> treatment <click on> treatments and side effects <click on> physical side effects <click on> fatigue.
- http://www.youtube.com/watch?v=YTFPMYGGe86s&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
- American Cancer Society: www.cancer.org <click on> treatment <click on> treatments and side effects <click on> physical side effects <click on> sexual side effects in women

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
Follow up Colon Cancer Care Schedule
You can use these charts to keep track of your follow up care.

<table>
<thead>
<tr>
<th>Family Doctor/Nurse Practitioner/Health Care Provider’s Visit</th>
<th>Year 1</th>
<th>Year 2</th>
<th>Year 3</th>
<th>Year 4</th>
<th>Year 5</th>
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<tr>
<th>CEA test at each doctor’s visit</th>
<th>Year 1</th>
<th>Year 2</th>
<th>Year 3</th>
<th>Year 4</th>
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<tr>
<th>CT Scan – Chest, Abdomen, Pelvis annually in Years 1-3. After Year 3, as your doctor feels it is needed.</th>
<th>Year 1</th>
<th>Year 2</th>
<th>Year 3</th>
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<th>Colonoscopy at 1 year after your diagnosis. If clear, it should be done at 3 years, 5 years and every 3-5 years after that as long as you are in good health.</th>
<th>Year 1</th>
<th>Year 2</th>
<th>Year 3</th>
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This booklet was approved by
Nova Scotia Cancer Patient Education Committee, 2014
This booklet is to be updated every three years


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Follow up Care for Colon Cancer Patients