Your Transplant Doctor is: _____________________________
Phone Number: _____________________________

Your Transplant Coordinator is: _____________________________
Phone Number: _____________________________

Your Transplant Social Worker is: _____________________________
Phone Number: _____________________________

Your Transplant Financial Coordinator is: _____________________________
Phone Number: _____________________________

Your Transplant NMDP Coordinator is: _____________________________
Phone Number: _____________________________

TABLE OF CONTENTS

Welcome ......................................................... 2
Allogeneic Blood and Marrow Transplantation: FAQ .......................... 3
Your Transplant Team .......................................... 4
Transplantation: Step By Step .................................. 6
Caregivers ....................................................... 8
Allogeneic Transplant Donors .................................. 9
Your Transplant Evaluation .................................... 10
Your Hospital Stay ............................................. 11
Hospital Transplant Admission ................................. 12
Transplant Day: Infusion of Stem Cells ....................... 14
Post-Transplant: What to Expect .............................. 15
Graft Versus Host Disease .................................... 17
Transition From Hospital to Home .......................... 18
Outpatient Follow-Up ......................................... 20
Guidelines at Home .......................................... 21
Nutrition ......................................................... 24
Long-Term Follow-Up ........................................ 28
Online and In-Person Support ............................... 28
Financial Resources ........................................... 30
Transportation .................................................. 30
Local Housing .................................................. 31
Recommended Books ........................................... 31
Welcome to Sutter Health Cellular Therapy Program

Considering an allogeneic stem cell transplant can be a frightening experience. You’ll likely have periods of mixed emotions, including both eagerness and anxiety about the process. You and your family will have many questions.

This guide provides information regarding allogeneic stem cell transplant, including the time before and after the transplant, the blood or marrow transplant (BMT) itself and how we’ll support you throughout the process.

The BMT staff shares a commitment to quality in medical and nursing care, lab services, rehabilitative therapy, social service support and clinical research to provide access to the latest treatments and therapies. Our team always keeps in close communication with your primary care doctor.

Keep this guide with you to review with your care team members and as new questions come up. There’s a lot to learn and you don’t need to learn it all at once. Pace yourself and review this handbook when needed. Your transplant team is available to answer any questions you have.

Allogeneic Blood and Marrow Transplantation: FAQ

What are stem cells?

Stem cells are “baby” blood cells, or blood-forming cells, produced in your bone marrow. These stem cells eventually become adult blood cells, particularly red blood cells, platelets and white blood cells. Each of these cells has a very specific task:

- **Red blood cells** carry hemoglobin, which supplies oxygen throughout your body to tissues and organs.
- **Platelets** clot the blood and prevent bleeding.
- **White blood cells** fight infection.

What is a blood or marrow stem cell transplant?

The term **stem cell transplant** describes a transplant using bone marrow or stem cells found in the blood. A blood or marrow stem cell transplant provides a possible cure for blood cancer, or a longer period of disease-free survival. The transplant starts with chemotherapy, radiation or both to deplete the cancer or diseased blood cells. The patient then receives an infusion of healthy cells into the bloodstream.

In a bone marrow transplant, stem cells are taken from the marrow of the donor or patient. In a peripheral blood stem cell transplant (PBSCT), stem cells that circulate in the bloodstream are collected through a venous access. Healthy stem cells are then infused into the venous access device in your vein.

What does allogeneic mean?

The root “allo” means “other.” With an allogeneic transplant, you receive healthy stem cells from a source other than yourself, such as a matched donor (related or unrelated) or donated umbilical cord blood, to replace your own damaged or diseased cells.

Why are stem cell transplants performed?

A stem cell transplant is used when:

- Your body’s bone marrow cannot or fails to make the blood cells your body needs.
- Disease compromises your bone marrow or blood cells.
- You’ve received high doses of chemotherapy or radiation, and your body needs a replacement of cells lost during these treatments.
- Your body’s stem cells don’t develop into the blood and other components your immune system needs to function.
- Your cancer returns after previous treatment.
- Transplant is the best treatment for disease control.
Your Transplant Team

A blood or marrow transplant is more than just a series of medical procedures. It’s a challenging experience for you, your family and other loved ones. That’s why we gather a team of skilled professionals to help meet your needs. Our goal is to create a partnership of care with you and your family.

These are some of the team members that may contribute to your care.

BMT RN Coordinator

You’ll be assigned a BMT RN coordinator early in the pretransplant phase. The coordinator is a registered nurse who specializes in stem cell transplants. Their primary role is to teach both you and your caregiver about the stem cell transplant before, during and after the procedure and help guide and support you and your care team throughout the transplant process.

National Marrow Donor Program (NMDP) Coordinator

A NMDP coordinator provides support throughout the process. The coordinator works with you and your care team to coordinate and organize the search for a matched donor or cord blood unit through the National Marrow Donor Program (NMDP) if a related donor is not available. The coordinator contacts all national and international unrelated donor registries and cord blood banks as part of the search for your stem cell donor.

Once a donor is found, the NMDP coordinator partners with your care team, the donor registry and donor center to schedule the stem cell harvest or collection at the appropriate time and under the direction of your BMT doctor.

Social Worker

This process is an emotional experience. To support your mental health during all stages of your transplant, a social worker will meet with you and your family to assess any barriers to care and be sure you’re managing well. The social worker will:

• Help you establish housing in the area, if needed, and create a caregiving plan.
• Perform a required mental health assessment as part of the insurance authorization process.
• Help you complete an advanced care plan, which may include a POLST (Physician Orders for Life-Sustaining Treatment). This gives your doctor specific instructions about what treatments you’d like to have at the end of life.
• Assist you and your caregiver with the Family Medical Leave Act (FMLA) or Paid Family Leave (PFL) process.

Financial Coordinator

A financial coordinator’s role is to help you understand the cost of a transplant and what’s covered by your insurance. They will work with you to provide guidance between the hospital and your insurance company. Call the financial coordinator with any questions or concerns about your potential financial needs.

Bedside Transplant Nurse

Being in the hospital can be overwhelming. Your bedside nurse is specially trained and knowledgeable to guide and care for you while you’re in the hospital. Your bedside transplant nurse collaborates with the transplant doctor and the rest of the transplant team to ensure your needs are met.

Case Manager

As part of the inpatient team, a case manager helps anticipate needs in advance of discharge to help ensure a smooth transition home.

Pharmacist

The pharmacist reviews all medication and treatment orders written by your doctor. Your pharmacist is available to answer any medication questions you might have.

Radiation Oncologist

If your treatment plan includes radiation therapy, you’ll meet with a radiation oncologist to discuss it.

Infusion Center Nurses

After discharge from the hospital, you’ll be returning to the infusion center. Infusion center nurses will assist you with any additional treatments after your hospital stay, such as lab draws and blood transfusions.

Clinical Dietitian

Healthy foods are an important part of recovery. A dietitian is available to meet with you before, during and after your transplant to assist you in meeting your nutritional needs. If your appetite isn’t good during your hospital stay, the dietitian will work with your doctor to determine the best way to get your body the nutrition it needs to heal. After transplant, if you have problems with eating or drinking, you may continue to see a dietitian as an outpatient.

Physical Therapist

Exercise is important before, during and after treatment. Someone from physical therapy will talk with you on admission. The physical therapist will design an individual exercise program to help you maintain strength and endurance during your hospital stay.

Chaplain

If helpful to you or your family, chaplains are available to talk about your spiritual needs. Chaplains are nondenominational and not affiliated with a specific religion.

Music Therapist

Music can be very healing. A certified music therapist may visit you at the bedside and play music for you. If you’re up for it and want to play music yourself, the music therapist has access to different instruments they can leave with you. Some patients even write songs or learn to play the basics of an instrument while in hospital.

Massage Therapist

Like music, touch can be very therapeutic. A licensed massage therapist may offer relaxation techniques to help you work through any anxiety or pain while you’re in the hospital.
Step 1: Planning Ahead
- Meet your transplant team.
- Assess financial coverage and other financial resources.
- Assess options for sperm banking (males) and egg or embryo freezing (females).
- Meet with BMT social worker.
- Complete financial applications (see page 30). Social worker to assist.
- Select caregiver(s).
- Make decisions about childcare and schooling.
- Arrange transportation to and housing near your center if needed (see pages 30-31). Decide on arrival date with assistance from BMT RN coordinator.
- Complete Advance Directives and POLST (social worker to assist).
- Decide if you need special preparations: dental, nutrition, fertility.
- For allogeneic transplant: complete human leukocyte antigen (HLA) typing for self and potential sibling donor(s).

Step 2: Preparation for Transplant
- Arrive to your transplant center.
- Meet with the BMT RN Coordinator for education.
- Complete medical evaluation with BMT team.
- After completing evaluation studies and obtaining insurance authorization, schedule an appointment to sign consent forms.
- For allogeneic patients with sibling donors: sibling donors will have tests to see if they’re a match and, if so, will need to sign consents.
- Receive placement of a central venous catheter.
- Stay active by walking or other mild exercise as suggested by your doctor.
- Ask questions and talk with staff regarding any concerns.

Step 3: Preparative Regimen (Conditioning)
- Admit to hospital.
- Receive pretransplant chemotherapy through your central venous catheter.
- Donor cells are collected from your donor.
- Activities: frequent mouth rinses, daily showers, walking three times per day.
- Take medication to help prevent symptoms and keep you comfortable.
- Let staff know how you’re doing and what you need.

Step 4: Transplant
- It’s a big day! Infusion of stem cells.

Step 5: Waiting for Engraftment
- Nursing staff monitors your blood counts daily.
- Receive transfusions of red blood cells and/or platelets if needed.
- Tell your doctor and nurse about any side effects: nausea, diarrhea, taste alteration, sores in mouth or anything else that isn’t normal for you.
- Talk with the support services team for your mental health. The team includes a nondenominational chaplain for spiritual needs, a social worker for problem solving and counseling, a physical therapist and a dietitian.
- Stay active. Talk with a physical therapist about exercises you can do in your room.
- Set goals for each day and cheer yourself on when you meet them.

Step 6: Recovery
- Discharge from hospital: time to go home!
- Your caregiver will be responsible for helping you manage your medicines, watching out for symptoms, driving you to doctor appointments and more.
- After you go home, you’ll have frequent or daily appointments in the infusion center, including blood draws and transfusions.
- Focus on food. Encourage yourself to eat the foods your dietitian suggests.
- Increase your activity. Talk with a physical therapist about how you can get stronger.
- Relax and nap during the day. Rest is important.
- Take things one day at a time and keep track of your daily goals to see your progress.
- Follow up with your BMT doctor at 30, 100 and 180 days after transplant, one year after transplant, and then annually for evaluation.
Caregivers

A caregiver is a family member or friend who can help you before, during, and after your transplant. Your caregiver will provide emotional support, care for you when you’re not feeling well and work with your team to report any side effects or concerns you may have before, during or after treatment.

Prior to starting the transplant process, you’re required to identify at least one person to be your caregiver. Your caregiver will need to be available 24 hours a day, seven days a week for general care, emergencies and to drive you to and from appointments. You may also choose two to three backup caregivers who may take turns staying with you. Due to your increased risk, your caregiver(s) must agree not to receive any live-virus vaccinations during and after your transplant for 100 days.

Caregiver Role Before Transplant

1. Participate in the pretransplant education meeting(s) with the BMT RN coordinator.
2. Participate in the consent signing process and sign the caregiver contract.
3. Accompany you to doctor and evaluation appointments as needed. This will include the day of central line placement as well as chemotherapy administration. (You’ll receive a calendar with these dates.)
4. Provide 24/7 observation and support.
5. Provide or arrange transportation to and from the clinic and/or infusion center daily.
7. Keep track of symptoms daily.
8. Record and refill medications.
9. Assist you with taking medications on schedule.

Caregiver Role After Transplant

1. Be available 24/7 to assist with post-transplant recovery.
2. Encourage fluids and food as tolerated.
3. Live or stay within 20 minutes of the hospital.
4. Provide or arrange transportation to and from the clinic and/or infusion center daily.
5. Take notes during appointments.
6. Perform housecleaning before and after discharge.
7. Meal management: prepare meals and snacks, encourage eating, meal cleanup.
8. Monitor symptoms daily with the Stoplight Tool you will receive from the coordinator. Take the patient’s temperature twice a day, or more often if needed, and record. Notify doctor if temperature is higher than 100.4°F, and/or you experience diarrhea, nausea, vomiting or dehydration.
10. Take you to the emergency room or hospital if needed.

You may choose a few backup caregivers who take turns staying with you.

Allogeneic Transplant Donors

With an allogeneic transplant, you receive stem cells collected from a donor who can be related or unrelated.

HLA Typing

Human leukocyte antigen (HLA) typing is a special blood test to determine your white blood cells’ unique “fingerprints,” or antigens. These antigens are inherited from parents—half from a mother and half from a father. You’ll undergo HLA typing once your transplant doctor determines that allogeneic transplant is the best treatment option for you.

Related Donor

Your transplant doctor will gather information, including HLA typing, from your first-degree relatives—siblings, parents and/or children. Results may take at least two full weeks for each full sibling. A patient has a one in four (25%) chance of a full match. Types of related-donor transplant include:

- Matched sibling transplant – Cells from an identical HLA-matched sibling.
- Haploidentical transplant – Cells from a patient’s parent or child.
- Syngeneic transplant – Cells from an identical twin.

Only about 30% of patients have an acceptable HLA-matched donor within their family, so an unrelated donor search may be needed.

Unrelated Donor

If there’s no match within your family, your transplant team will work with the National Marrow Donor Program (NMDP) to find a donor for you.* It may take several months to find the best possible unrelated donor. While waiting for an unrelated donor, you may continue to receive additional treatment.

Once your team identifies potential donors, they’ll recommend which ones should proceed to a formal donor search. During this search, they’ll ask donors who are identified as the best possible matches to go to the nearest donor center for additional testing.

Matched volunteer donor types include:

- Matched unrelated donor – Cells collected from an HLA-matched volunteer donor obtained through NMDP.
- Cord blood transplant (CBT) – Cells from one or more HLA-matched umbilical cord(s) obtained through the NMDP.

While going through tests and evaluations, your donor will prepare for stem cell donation. After the donor passes the medical exams and signs all consent and other forms, they’re ready to donate their cells and a transplant date will be scheduled.

Stem Cell Mobilization

Mobilization is a process to increase blood stem cells in the bone marrow and force them into the bloodstream to be collected for transplant. Donors will receive injections to stimulate stem cell production and help move stem cells from the bone marrow to the blood. This will take place before your admission to the hospital.

Stem Cell Collection Methods

Once your donor’s stem cells are ready, they can be collected in one of three ways:

- Apheresis – This procedure removes stem cells from the blood and usually takes two (2) eight-hour days.
- Bone Marrow Harvest – This collects cells directly from bone marrow in the hip during an operating-room procedure.
- Cord Blood Products – Blood from a baby’s umbilical cord, stored in a blood bank, is sometimes used for stem cells. Cord blood products are obtained through the National Marrow Donor Program.

*Contact With Your Donor: The transplant team can’t share any information about potential donors until the donor has been selected. At that point, you can learn only the donor’s age and gender. One year after transplant, you and your donor may write letters anonymously through the NMDP. If both you and the donor are willing, you may learn each other’s identity through the NMDP.
Prior to transplant, you’ll undergo several tests, evaluations and procedures to ensure you’re in good physical condition. These may include:

- **Central venous catheter.** A surgeon or radiologist will insert a catheter – a small flexible tube – into a large vein in your chest, just above your heart. The catheter provides a route to administer all chemotherapy drugs, blood products and other intravenous fluids. It’s also used to withdraw blood for the many samples needed throughout your hospital stay.

- **Chest X-Ray.** A chest X-ray to evaluate your lungs.

- **Cytomegalovirus (CMV) Infection Test.** Cytomegalovirus (CMV) belongs to the herpes family of viruses, which includes the viruses that cause chicken-pox, shingles and cold sores, among others. It’s a common infection, and many adults are CMV positive. CMV positive means that you have been exposed to the virus in the past and it may still exist in your body without having an active infection or you even feeling ill.

CMV usually stays hidden in your body unless your immune system starts to weaken, like it does during a stem cell transplant. If this happens, the virus can attack the lungs, gut, liver or nerves, and in severe cases, can be fatal.

Prior to transplant, you’ll be tested to determine your CMV status. If you are CMV negative, great effort is made to keep you, the patient, negative. This includes giving you blood and stem cells from those who are also CMV negative, if possible.

**Your Hospital Stay**

If you’re being admitted to the hospital for transplant, follow these guidelines to ensure your safety, promote a healing environment and support a quality health-care experience.

**Blood and Marrow Transplant (BMT) Unit Guidelines**

The BMT unit is a positive-pressure unit with HEPA air filters. Vistation is limited to two guests at a time. Hospital visiting guidelines may change according to state mandates and other circumstances. Children may visit when accompanied by an adult.

No guests may enter the BMT unit if they have symptoms of illness, such as cough, fever, running nose, malaise, nausea or vomiting. Everyone must wash their hands before entering and when leaving the BMT unit and patient rooms. Hand sanitizer and handwashing sinks are readily available. Latex balloons, fresh flowers, plants and pets are not allowed.

One visitor over age 18 may stay overnight in your room. Visitors may not use patient restrooms. Visitor restrooms are in the family waiting room.

**Expectations During Your Stay**

- **Social work assessment.** This evaluation helps your team better understand your readiness and understanding of the transplant and the support you may need.

- **Dental exam.** If you haven’t seen your dentist in the past six to nine months, you’ll need an exam and/or dental clearance. Simple cavities may be filled, but if you need any major work, you’ll need to get approval from your transplant physician. You may need antibiotics prior to any dental work, or you may need to wait until well after the transplant to have dental work done.

- **Electrocardiogram (EKG).** This test examines the heart’s electrical activity.

- **Multigated acquisition (MUGA).** This test measures how well your heart pumps blood through your body.

- **PET/CT or CT scan.** A full-body scan looks for any additional disease in other areas.

- **Pulmonary function test (PFT).** This test examines how well your lungs are working. You’ll also have blood drawn from your wrist to measure arterial blood gas (ABG).

- **Brush your teeth and/or use a mouth rinse at least four times a day, typically after meals and before bed. This reduces infection risk.**

- **You’ll receive nurse visits every hour to deliver medications, provide treatments and perform assessments. You’ll have a blood draw between 4-5 a.m. daily.**

- **The vascular access team checks your central line every day to assess its condition and prevent infection.**

- **The hematology-oncology team will visit daily. This is a good time to discuss your care and ask questions.**

**How To Promote Your Health and Comfort**

- **Bring comfortable shoes.** Walk the length of the BMT Unit three times a day to maintain muscle strength.

- **Bring clothing that allows easy access to your central line.** You can wear these clothes or a hospital gown.

- **Bring personal items such as books, computer, gaming devices or games to pass the time.** We are not responsible for items that may become lost or stolen.

- **The unit has a cart-mounted TV you can check out for personal gaming, and you can access the internet through SHGuestNet.**

- **Bring personal items such as books, computer, gaming devices or games to pass the time.**

- **A charge nurse is always available to manage any concerns or questions.**

- **Your medications may alter the taste of food.** Your family may bring in food or have it delivered, and you can talk with the dietitian for help with food selection. Please avoid raw or undercooked meats and unwashed fresh fruits and vegetables. Make sure that foods are freshly prepared and stored properly. All food stored in the family room refrigerator must be labeled with your name and the date.
How Friends and Family Can Reach You in the Hospital

Your room phone can receive direct calls from 7 a.m. to 10 p.m.

Because nurses are usually in patients’ rooms and unable to answer calls, family and friends should not call the nurses’ station but instead call your room directly, unless there’s a specific problem or issue that needs to be addressed with the staff.

We will not release any information about you to any caller without your permission. Please identify one family member or friend as the primary person to call for information. If you’d prefer not to receive calls in your room, simply let the nursing staff know to hold your calls.

Preparative Regimen (Conditioning)

Before you’re admitted to the hospital, usually during consent signing, your doctor and BMT coordinator will discuss your particular preparative regimen (conditioning) with you and answer any questions. If you have additional questions during the regimen, please ask your nursing staff.

Pretransplant preparation destroys healthy and unhealthy cells. This process includes chemotherapy with or without total body radiation and is based on a specific treatment protocol designed for you. The length of time may vary from one to eight days before transplant.

Shortly after arrival in the unit, you’ll begin your preparative regimen. Your central venous line will be placed, you’ll start receiving intravenous (IV) fluids and your chemotherapy will be ordered from pharmacy.

Registration and Arrival at the BMT Unit

On admission day, please arrive by 7:30 a.m.

Register at the desk in the lobby of the Anderson Lucchetti Women’s and Children’s Center on the main campus. From there, you’ll be escorted to the 4th floor BMT Unit. Upon admission to the transplant center, the nursing staff will take your vital signs and orient you to your room and the unit. Your transplant doctor will visit to ensure that you’re ready to proceed.

While in the hospital, your nursing staff will keep you informed about your care and provide hourly safety rounds. At the beginning of each shift, your nursing team will discuss your care during a bedside report. We encourage you to bring up any questions or concerns.

Transplant Medications

Reactions vary when receiving chemotherapy. The most common complaints are nausea with or without vomiting, plus fatigue. Other side effects may include diarrhea, loss of appetite, mouth sores, hair loss and infertility.

Medications can address many of these side effects. You may be taking these medications during your stay and when you go home. Since the drugs work without vomiting, plus fatigue. Other side effects may include diarrhea, loss of appetite, mouth sores, hair loss and infertility.

Medications can address many of these side effects. You may be taking these medications during your stay and when you go home. Since the drugs work without vomiting, plus fatigue. Other side effects may include diarrhea, loss of appetite, mouth sores, hair loss and infertility.

In addition to chemotherapy or fluids, medications used during your stay may include:
- Antibiotics to prevent or treat infections
- Antinausea medicine to prevent or treat nausea
- Antifungals to prevent or treat fungal infections
- Antivirals to prevent or treat viral infections
- Narcotics to relieve pain
- Electrolyte replacement

While you’re in the hospital, you’ll receive immunosuppressants through your IV. Your medical team will watch the levels closely and may adjust the dose.

Once you go home, you’ll receive immunosuppressants in pill form. You’ll have a daily blood draw at the infusion center to monitor the medication. Don’t take your morning dose, but bring your pills with you each day so you can take them immediately after you have your blood drawn.

The evening dose may be different from your morning dose, depending on your lab results. The clinic will notify you of any changes to the dose.

Ask your medical team if you have any questions regarding your medications. Before your hospital discharge, you must have all your medications on hand; your medical team will provide a list and review each one with you and your caregiver. It’s important to take your medications as directed by your doctor.

Immunesuppressants (Antirejection Drugs)

This important type of medication helps prevent graft-versus-host disease (GVHD). As the name suggests, immunosuppressants suppress your immune system so it won’t attack your donor’s cells.

While you’re in the hospital, you’ll receive immunosuppressants through your IV. Your medical team will watch the levels closely and may adjust the dose.

Once you go home, you’ll receive immunosuppressants in pill form. You’ll have a daily blood draw at the infusion center to monitor the medication. Don’t take your morning dose, but bring your pills with you each day so you can take them immediately after you have your blood drawn.

The evening dose may be different from your morning dose, depending on your lab results. The clinic will notify you of any changes to the dose.

Ask your medical team if you have any questions regarding your medications. Before your hospital discharge, you must have all your medications on hand; your medical team will provide a list and review each one with you and your caregiver. It’s important to take your medications as directed by your doctor.

Other Medications

• Electrolyte replacement
Transplant Day: Infusion of Stem Cells

Stem cell infusion is similar to blood transfusion. The total time may vary, depending on the volume of stem cells, how well your body tolerates the infusion and whether the cells are fresh or frozen.

At least two nurses will be with you before and during the infusion to monitor your vital signs. Your transplant doctor will also be available during the infusion. If you have any questions or concerns, please ask your nursing staff.

Frozen Stem Cells

Cells may have been collected, preserved with the chemical dimethyl sulfoxide (DMSO) and delivered to the transplant unit frozen. Nurses thaw the frozen cells in a lukewarm water bath. Once thawed, the cells can be infused like any other blood product.

DMSO enters your bloodstream during the infusion. Your body removes it through breathing, so take deep breaths in and out to help this process along. Patients commonly report that DMSO has a strong, garlic-type taste, while others report a sweet corn taste. You might want to have hard candy (preferably on a stick) at your bedside during the infusion. It’s also common for you, your room and the unit to smell like garlic for a while after the infusion.

Fresh Stem Cells

Allogeneic donor stem cells may be collected the day before transplant and are infused as fresh cells, much like a blood transfusion.

Possible Side Effects

You’ll receive medicine prior to the infusion to help reduce possible side effects, such as:

- Nausea
- Vomiting
- Fever or chills
- Shortness of breath or change in oxygen level
- Low or high blood pressure
- Increased heart rate
- Rash or hives
- Discolored or reddish urine

You’ll see a special supplies cart placed outside or near your room or chair.

Post-Transplant: What To Expect

For many people, the most difficult time of the transplant process is between infusion and discharge.

Engraftment

Approximately 10-28 days after transplant, your medical team expects to see signs that new bone marrow or stem cells are growing (engrafting) and beginning to produce blood cells. The medical team usually monitors blood counts daily. A rising white blood cell count is often the sign of a successful transplant, or graft. You may want to keep a daily record of your blood counts to track your progress.

In addition to watching for the first signs that your new immune system is working, your medical team will monitor you closely to detect any complications. If complications arise, treatment begins promptly.

Until the new cells engraft, you’re very susceptible to infections. During this time, your doctor may give you antibiotics. You may experience fever and will be monitored closely for any symptoms of infection. Because your hemoglobin may be low, you may feel tired. Because your platelets may be low, you may experience bleeding; if so, please notify your nurse and doctor immediately. Your medical team will monitor all your lab values closely and you’ll be given either red blood cells or platelets as needed. Despite fatigue, it’s important to maintain activity to prevent complications.

Maintaining Your Well-Being

During transplant and after, be prepared for the possibility of an emotional roller coaster. You may feel:

- Vulnerable and anxious while you’re waiting for engraftment and not protected by a fully functioning immune system.
- Intense concerns about physical symptoms.
- Worry about making it through transplant.
- A loss of personal control as you cope with the daily monitoring and medical procedures.

All of these feelings are common but coping with them can be a challenge. It may help to talk with others, keep active and engage in activities that help take your mind off treatments. You may find that working out the details of your day, such as time of bathing, walks and treatments, helps you maintain control and give structure to this period.

Be prepared for the possibility of an emotional roller coaster.
Potential Side Effects

It’s important to understand the risks. You may or may not experience any of these, but be aware of what to look for and what to report.

- **Bleeding** – The transplant process stops the body’s platelet production. Platelets are essential to prevent bleeding and aid in the clotting process. The staff will monitor your platelet count daily and administer a platelet transfusion when needed. They’ll also monitor your urine, stool and any vomit for early signs of bleeding. To prevent bleeding from injury and for your safety, always call your nurse for help when you want to get out of bed.

- **Cataracts** – Your eye lenses may become cloudy after your transplant. If left untreated, this can cause partial or complete blindness. Establish regular care with an eye doctor before, during and after transplant.

- **Early Menopause** – Many women go into early menopause after transplant because the chemotherapy or radiation can cause the ovaries to stop producing eggs. This is especially likely if you’ve been on estrogen therapy. Side effects may include hot flashes, night sweats or irritability. You’ll also be at a higher risk for osteoporosis (bone thinning).

- **Graft Failure** – There’s a very rare (less than 1%) possibility that your stem cells won’t grow back to a fully functioning bone marrow. This can affect all the blood cells or only specific types. You’ll receive lab tests daily to monitor engraftment.

- **Hemorrhagic Cystitis** – Your bladder lining may become irritated and bleed. Symptoms are similar to a urinary tract infection. You’ll receive large volumes of IV fluids to protect your bladder. The nurses will also monitor how much urine you produce and will check urine for blood. In severe cases your doctor may ask for a bladder catheter to help keep you comfortable.

- **Infection** – Because your immune system is compromised, infections can develop. Your medical team will monitor your temperature and other vital signs frequently, even through the night. If you develop signs of an infection, you’ll receive additional medications to treat it. It’s possible that an infection may threaten to overwhelm your system, a condition called sepsis or sepsisemia that can be life threatening.

- **Lung Complications** – The lungs are a common infection site in blood marrow transplant patients, in part because most bone and marrow patients spend a great deal of time in bed. While full-blown pneumonia can be life threatening, it can be prevented. The nurses and the physical therapists will constantly encourage you to get out of bed, sit in a chair, walk in the hallway, exercise, breathe deeply and use an incentive spirometer (a breathing exerciser). Your own hard work can help prevent a serious complication.

- **Secondary Cancers** – Transplant patients are at higher risk for secondary cancer – usually leukemia (cancer of the blood cells). This can be related to chemotherapy or radiation treatment.

- **Short-Term Memory Loss** – Many transplant patients complain of short-term memory loss, which may be temporary or permanent. Some people find it helpful to keep a small notebook.

- **Sterility** – Your ability to have children may be affected by your treatment and transplant. Please refer to page 6 for information about fertility preservation.

- **Veno-Occlusive Disease** – Chemotherapy or radiation therapy may affect your liver. Your belly may become tender and swollen, and your skin and eyes may turn yellow. This condition usually heals on its own, however, severe cases can be life threatening. To monitor this, you’ll be weighed twice a day while in the hospital.

You may not experience any of these risks, but be aware of what to look for.

Graft Versus Host Disease (GVHD)

Graft versus host disease (GVHD) is a frequent complication of allogeneic transplants. GVHD means the donor’s cells attack your organs and tissues. Symptoms vary, and they can be mild or life threatening. If you experience these or other symptoms, please notify your transplant doctor and care team.

In general, 35% of allogeneic transplant patients develop some acute GVHD and 50% develop chronic GVHD.

**Acute GVHD**

Acute GVHD typically occurs during the first three months following an allogeneic transplant. It primarily affects the skin, liver, stomach or intestines. Doctors grade the severity of acute GVHD according to the number of organs involved and the degree to which they’re affected. Acute GVHD may be mild, moderate, severe or life threatening. Stages of acute GVHD are:

- **Stage 1 (mild):** a skin rash over less than 25% of the body.
- **Stage 2 (moderate):** a skin rash over more than 25% of the body accompanied by mild liver or stomach and intestinal disorders.
- **Stage 3 (severe):** skin redness similar to a severe sunburn, and moderate liver, stomach and intestinal problems.
- **Stage 4 (life-threatening):** blistering, peeling skin, and severe liver, stomach and intestinal problems.

**Chronic GVHD**

Chronic GVHD can develop after the third month following an allogeneic transplant and may persist over time. You may experience symptoms or side effects such as:

- Skin problems: dry itching rash, lesions, change in skin color or skin tightening.
- Partial hair loss or premature graying.
- Liver abnormalities: jaundice (yellowing of skin or eyes) and abnormal liver test results.
- Eye dryness or stinging.
- Difficulty with swallowing and eating.
- Burning sensation in the mouth when using toothpaste or eating acidic foods.
- Digestive problems: heartburn, stomach pain, weight loss, reduced nutrient absorption.
- Contractures: tightened tendons, difficulty extending or bending arms and legs.
- Lung problems: wheezing, bronchitis or pneumonia.
- Painful intercourse (women)

**Most Common Chronic GVHD Symptoms**

- Rash, itching, skin redness
- Dark spots, skin tightness, skin lesions
- Abnormal liver tests, jaundice
- Dry, burning eyes
- Dryness or sores in mouth
- Difficulty swallowing
- Difficulty breathing
- Contractures
- Pain with sexual intercourse

**Less Common Chronic GVHD Symptoms**

- Skin scarring
- Partial hair loss, premature graying
- Severe liver disease
- Vision impairment
- Heartburn, stomach pain
- Weight loss
- Bacterial infection, bronchitis, pneumonia
Transition from Hospital to Home

Discharge
You may feel a wide range of emotions as you think about going home. While you look forward to returning home:
• You may feel anxious about leaving the protected environment of the hospital and the security of the staff.
• You may be worried about caring for yourself at home.
• Your family or caregiver might have concerns about their ability to handle responsibilities that were once those of the hospital staff.
• You may wonder how other people will relate to your physical and other changes. Some people may not understand what you’ve gone through and how it has changed you, and you may miss the support system you had in the hospital.
• You’ll probably feel frustrated or down at times because of your isolation and limited activity.

These feelings are normal, and you may wish to seek out a supportive friend or relative who’s able to provide the understanding you need. The blood and marrow transplant team can also help. Above all, as you make this transition from hospital to home, remember to be patient, pace yourself and set small goals for yourself and others.

Medications
As you near your discharge date, your doctor and nursing staff will order your discharge medications. Your caregiver will need to pick up your medications and bring them to the hospital ahead of discharge to give your care team time to provide medication education.

When you’re discharged from the hospital, you’ll have thorough medication instructions, including a written schedule, and all of your medications in hand. Take your medications as directed and bring your medications with you to each infusion center visit.

Home Environment
Before leaving the hospital, your caregiver should ensure that your recovery location has been thoroughly cleaned. Someone other than you will need to be responsible for keeping the house picked up, especially the bathroom and kitchen. The goal is to reduce your exposure to germs while your immune system recovers.

While it’s important to keep the house clean, it shouldn’t take more than a few minutes a day. A caregiver needs to:
• Vacuum every day.
• Wipe down kitchen counters with a disinfectant or bleach cleaner daily.
• Use dish brushes (not dishrags) and wash them frequently.

You, the patient, should not change cat litters, pick up pet poop, dig in soil, tend to farm animals or engage in house renovations.

Your home needs to be a safe place for you. Don’t visit with anyone who has illness symptoms or who has been near anyone who’s ill.

Immediately after discharge, you’ll need to stay within a 20-minute drive to the hospital.
Outpatient Follow-Up

Doctor Visits
After you’re discharged from the hospital, you’ll remain under your transplant doctor’s care for a while. You’ll visit with the doctor approximately two to three times each week for a couple of weeks. During these appointments, ask questions or discuss concerns with your care team.

Contact your transplant team if you experience:
• Any type of bleeding.
• Burning, pain, increased frequency or urgency with urination.
• Discharge or odor from the vagina or penis.
• Headache or neck pain.
• Mouth sores, sore throat or trouble swallowing.
• New or persistent cough.
• New or persistent diarrhea or constipation.
• Physical changes or any other differences that concern you.
• Rectal soreness or abscesses.
• Redness or swelling on your skin.
• Redness, swelling or drainage at your catheter site.
• Shortness of breath while at rest or when exercising.
• Temperature of 100.4°F or higher.

If you need to go to an emergency department (ED), please contact your transplant team so you’ll be treated as a post-transplant patient and possibly admitted to the BMT Unit.

Infusion Center
After discharge, you’ll make daily visits to the infusion center for a period of time. The visits usually last one to two hours but may be as long as six hours. Your team will schedule your first infusion-center appointment before your hospital discharge. At your visits, infusion center nurses will check your lab values, check your vital signs and administer IV fluids, medications or blood products as needed.

Caring for Your Central Venous Catheter*
If your central infusion line gets damaged, it won’t work correctly and could raise your chance of getting an infection. To protect the central line at home:
• Use good hand hygiene. (see Hand Hygiene, page 21). Don’t touch the catheter or dressing. Always clean your hands before and after you touch any part of the central line. Your caregivers, family members and any visitors should use good hand hygiene, too.
• Avoid damage. Don’t use any sharp or pointy objects around the catheter. This includes scissors, pins, knives, razors or anything else that could cut or puncture it. Don’t let anything pull or rub on the catheter, such as clothing.

Call your healthcare provider right away if:
• Your dressing is damaged, dirty, wet or peeling off.
• You have redness, swelling, tenderness, leakage, bleeding or drainage around your catheter exit site.
• Your needleless connector falls off.
• You have chills or a fever of 100.4°F (38°C) or higher.
• You have a break or leak in your catheter.

* See page 10 for diagram

Guidelines at Home

Hand Hygiene
Just as when you were in the hospital, you must follow certain procedures at home to help prevent infection. Wash your hands frequently, and encourage visitors and family members to do so, particularly:
• Before and after using the bathroom.
• Prior to eating.
• Before and after preparing food.
• After coming home from a public place.

To clean hands, follow these steps:
• Wet hands with clean, running water (warm or cold). Apply soap.
• Lather hands by rubbing them together with the soap. Lather the backs of your hands, between your fingers and under your nails.
• Scrub hands for at least 20 seconds. Need a timer? Hum the “Happy Birthday” song from beginning to end twice.
• Rinse hands well under clean, running water.
• Dry hands using a clean paper towel, or air dry them.

When you can’t use soap and water, use an alcohol-based hand sanitizer that contains at least 60% alcohol.

Showering
You can shower with your catheter in place. Do not submerge your catheter in water, such as a bath, pool or hot tub.

Each time you shower, cover your dressing completely with plastic wrap and tape or a new, single-use waterproof cover, such as Aquaguard®. You can purchase these online. To use the waterproof cover:
• Peel off the top and side strips.
• Place the top edge above your dressing. Don’t let the tape touch your dressing. Smooth the cover down over your dressing.

Temperature Checks
Take your temperature twice a day at home. If you don’t have a working thermometer, please tell the BMT staff.
• Take your temperature at the same times each day.
• Anytime you feel sick or think you have a fever, check your temperature. If you notice that your temperature is increasing, check it again within 30-60 minutes.
• Don’t drink anything hot or cold for at least 15 minutes before taking your temperature.
• Always take your temperature before you take fever-reducing medications.
• If your temperature is 100.4°F or higher with or without fever-reducing medications, notify your team immediately.
• Do not use aspirin, ibuprofen, acetaminophen or products that contain them unless advised by your transplant doctor.
When you’re out in the sun, wear protective clothing and use sunscreen.

Bleeding Prevention

It’s normal to need platelet transfusions for a while after transplant. Notify your transplant doctor if you notice any bleeding, including bleeding on your mouth, gums, urine or stool.

To help prevent bleeding:
• Use a soft toothbrush or dental swabs.
• Wear a seatbelt with shoulder harness when in the car.
• Do not use rectal thermometers, rectal suppositories or enemas. Use stool softeners or dietary fiber to prevent constipation.
• Use an electric razor until platelet count is over 50,000 without transfusions.
• Don’t use dental floss until platelets are over 50,000 without transfusions.
• Avoid forcefully blowing your nose.
• Avoid contact sports and other activities that may cause bleeding or bruising.
• Use proper lubrication during sexual activity (see page 23).

Skin Care

Showering daily helps reduce potentially harmful bacteria that live on your skin. Keep your skin hydrated by avoiding hot showers and using unscented lotion daily.

Your skin will be very sensitive to sunlight and will burn easily. Avoid exposure to bright sunlight for at least six months after your transplant. When you’re out in the sun, wear protective clothing such as long sleeves, pants and a hat with a brim. Use sunscreen with an SPF of at least 15 on exposed skin.

It’s OK to wear makeup. Use new products and don’t share them. Chipped nail polish and artificial nail products encourage the growth of bacteria and fungus so we do not recommend their use.

Mouth Care

Continue with good mouth cleaning after meals and at bedtime. Get regular dental cleansings, but avoid dental work (fillings, bridges, etc.) for six months after your transplant. If you must get dental work, your doctor may recommend antibiotics.

Contact Lenses

Don’t wear contact lenses during your admission. Check with your doctor before resuming contact lens wearing.

Vaccines

Don’t get any vaccines unless ordered by your transplant doctor. Check with your doctor before visiting with adults or children who have recently received vaccines.

Physical Exercise

It’s common to feel fatigue after an allogeneic transplant. Physical activity plays a vital role in your recovery. Most of your waking time should be spent out of bed. Maintain your activity level by developing a regular exercise routine. This may be as simple as taking a short walk. Talk to your healthcare team about appropriate physical activities.

Sexual Activity

When you first come home from the hospital you might feel very tired and hesitant about intimacy with your partner. You can hug, cuddle, snuggle, kiss, touch each other and sleep in the same bed with your partner unless your partner is ill. As you start to exercise and eat better and your blood counts improve, your sexual desire should return. You may have gentle sexual intercourse when your platelet count is above 50,000 without the need for transfusions. Please discuss any questions with your transplant doctor. Your partner may also have questions.

Remember that your body is still adjusting to the effects of treatment. Practice good hygiene before and after sexual activities. Use proper lubrication during sexual intercourse to avoid irritation and possible bleeding or infection; a water-soluble, nonirritating lubricant can help. Urinate after intercourse to prevent urinary tract infections.

Although women may go into early menopause after transplant, that doesn’t always happen and shouldn’t be counted on for birth control. Safe sex practices, including condom use, are always encouraged to prevent sexually transmitted diseases. Speak to your physician about contraceptive options, pregnancy concerns or any other questions you may have.

Activities Away From Home

Schedule out-of-home activities at nonpeak hours; for example, shop early in the morning when most people are at work. Avoid areas like construction sites which may have mold, fungus or bacteria.

Return to Work or School

You may require more time before you’re able to resume work or school, especially if you require further treatment after transplantation. Discuss this with your transplant team.

Children

Children are wonderful, but they can be little germ carriers, especially if they attend school. Reduce visits from children when possible. If your home includes schoolchildren, ask the school to notify you if there are any outbreaks of chicken pox, mumps or measles. The transplant doctor might want you to keep your child out of school for a short period of time.

If you must change a soiled diaper, wear disposable gloves and always wash your hands afterward.

Animals

Animals can be therapeutic, but all animals carry some risks. Avoid any aggressive animal that may bite, scratch or cause a fall. Don’t handle any animal waste, including poop scooping or cleaning litter boxes, fish tanks and bird cages. Always wash your hands after any animal contact.

Readmission

You may need to be readmitted to the hospital if you’re experiencing fever, infection or inability to drink or eat. This isn’t uncommon and doesn’t necessarily mean you’re not progressing as you should.

Schedule out-of-home activities at nonpeak hours.
After leaving the hospital, you may discover that foods still taste different and your appetite hasn’t returned to normal. Maintain a well-balanced diet with adequate calories and protein to continue to help your bone marrow and immune systems recover. To prevent dehydration, try to drink at least eight to 10 glasses of fluid each day.

Most of the nausea and vomiting associated with your transplant should be over. If not, your transplant doctor can prescribe an antinausea medication. It might be easier for you to eat several small meals and snacks rather than three large meals.

Use these neutropenic diet recommendations, specific for people with weakened immune systems. Eat food products prior to their expiration dates. If you aren’t sure whether you should eat something, don’t.

“Shelf stable” refers to unopened canned, bottled or packaged foods that can be stored at room temperature before opening; the container may require refrigeration after opening.

### Nutrition

<table>
<thead>
<tr>
<th>RECOMMENDED</th>
<th>NOT RECOMMENDED</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Beverages</strong></td>
<td></td>
</tr>
<tr>
<td>• Tap water</td>
<td>• Hot tea</td>
</tr>
<tr>
<td>• Commercial bottled distilled spring water</td>
<td>• Malt tea</td>
</tr>
<tr>
<td>• All canned, bottled, powdered beverages</td>
<td>• Unpasteurized beer</td>
</tr>
<tr>
<td>• Instant and brewed coffee or tea</td>
<td>• Well water, unless tested yearly for safety</td>
</tr>
<tr>
<td>• Beer pasteurized after fermentation</td>
<td></td>
</tr>
<tr>
<td>• Brewed herbal teas using commercial packaged tea bags</td>
<td></td>
</tr>
<tr>
<td>• Liquid or powdered nutritional supplements and baby formula</td>
<td></td>
</tr>
<tr>
<td><strong>Breads, Grains, Cereals</strong></td>
<td></td>
</tr>
<tr>
<td>• All breads, bagels, rolls, crackers, muffins, sweet rolls, biscuits, pancakes, waffles and French toast</td>
<td>• Raw grain products</td>
</tr>
<tr>
<td>• All cooked and ready-to-eat cereals</td>
<td></td>
</tr>
<tr>
<td>• All chips, pretzels, popcorn, tortillas</td>
<td></td>
</tr>
<tr>
<td>• Cooked rice, pasta, beans and other grains</td>
<td></td>
</tr>
<tr>
<td><strong>Fruits and Fruit Juice</strong></td>
<td></td>
</tr>
<tr>
<td>• Any canned, stewed or frozen cooked fruit</td>
<td>• Unwashed raw fruit</td>
</tr>
<tr>
<td>• Well-washed raw fruit (except as noted)</td>
<td>• Unpasteurized fruit juices</td>
</tr>
<tr>
<td>• Dried fruit</td>
<td>• Fresh strawberries and other berries</td>
</tr>
<tr>
<td>• Pasteurized fruit juices</td>
<td>• Fresh grapes</td>
</tr>
<tr>
<td><strong>Vegetables and Vegetable Juice</strong></td>
<td></td>
</tr>
<tr>
<td>• All cooked frozen, canned or fresh vegetables</td>
<td>• Unwashed raw vegetables and herbs</td>
</tr>
<tr>
<td>• Well-washed raw vegetables</td>
<td>• Vegetable sprouts</td>
</tr>
<tr>
<td>• Fresh, well-washed herbs and dried herbs and spices</td>
<td></td>
</tr>
<tr>
<td>• Pasteurized vegetable juices</td>
<td></td>
</tr>
<tr>
<td>• Bottled and canned salsa, refrigerated after opening</td>
<td></td>
</tr>
<tr>
<td><strong>Meat and Meat Substitutes</strong></td>
<td></td>
</tr>
<tr>
<td>• All well-cooked or canned meats (beef, pork, lamb, ham, bacon, sausage, hot dogs), poultry, game, fish and shellfish</td>
<td>• Raw or undercooked meats</td>
</tr>
<tr>
<td>• Well-cooked eggs (white cooked firm with thickened yolk)</td>
<td>• Raw or undercooked poultry, game, fish (including, sushi, sashimi, ceviche, pickled fish, cold smoked salmon lox)</td>
</tr>
<tr>
<td>• Pasteurized egg substitutes</td>
<td>• Raw or undercooked tofu or eggs (and foods containing them)</td>
</tr>
<tr>
<td>• Commercially packaged sausami, bologna and other lunch meats</td>
<td>• Deli meats, unless further cooked</td>
</tr>
<tr>
<td>• Canned and commercially packaged hard-smoked fish, refrigerated after opening</td>
<td>• Fermented soybean products such as tempeh</td>
</tr>
<tr>
<td>• Cooked tofu</td>
<td></td>
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<tr>
<td><strong>Milk and Milk Products</strong></td>
<td></td>
</tr>
<tr>
<td>• All pasteurized Grade A milk, milk products, commercially packaged cheese and cheese products</td>
<td>• Unpasteurized or raw milk, unpasteurized cheese, yogurt and other milk products</td>
</tr>
<tr>
<td>• Pasteurized yogurt and sour cream</td>
<td>• Cheese with molds (blue stilton, Roquefort, gorgonzola)</td>
</tr>
<tr>
<td>• Dry, refrigerated and frozen pasteurized whipped topping</td>
<td></td>
</tr>
<tr>
<td><strong>Fats and Oils</strong></td>
<td></td>
</tr>
<tr>
<td>• Refrigerated commercial and homemade cakes, pies, pastries and puddings</td>
<td>• Unrefrigerated, cream-filled pasty products (not shelf stable)</td>
</tr>
<tr>
<td>• Refrigerated, cream-filled pastries</td>
<td></td>
</tr>
<tr>
<td>• Homemade and commercial cookies</td>
<td></td>
</tr>
<tr>
<td>• Ices, popsicles, gelatin desserts</td>
<td></td>
</tr>
<tr>
<td>• Candy, gum</td>
<td></td>
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<tr>
<td>• Shelf-stable, cream filled cupcakes, fruit pies and canned pudding</td>
<td></td>
</tr>
<tr>
<td>• Commercially packaged ice cream, frozen yogurt, sherbet, ice cream bars</td>
<td></td>
</tr>
<tr>
<td>• Homemade milkshakes</td>
<td></td>
</tr>
<tr>
<td><strong>Desserts</strong></td>
<td></td>
</tr>
<tr>
<td>• Cooked gravy and sauces</td>
<td></td>
</tr>
<tr>
<td>• Nondairy creamers</td>
<td></td>
</tr>
<tr>
<td>• Nondairy whipped topping</td>
<td></td>
</tr>
<tr>
<td><strong>Soups</strong></td>
<td></td>
</tr>
<tr>
<td>• All canned and homemade cooked soups</td>
<td>• All miso products (miso soup)</td>
</tr>
<tr>
<td><strong>Miscellaneous</strong></td>
<td></td>
</tr>
<tr>
<td>• Salt, sugar, brown sugar</td>
<td>• Raw or unpasteurized honey</td>
</tr>
<tr>
<td>• Jam, jelly and syrups, refrigerated after opening</td>
<td>• Herbal and nontraditional nutrient supplements</td>
</tr>
<tr>
<td>• Commercial (heat treated) honey</td>
<td>• Brewer’s yeast, if eaten uncooked (avoid contact with raw yeast)</td>
</tr>
<tr>
<td>• Ketchup, mustard, BBQ sauce, soy sauce, pickles, pickle relish, olives, and other condiments, refrigerated after opening</td>
<td></td>
</tr>
</tbody>
</table>
Food Handling Guidelines

People with a compromised immune system are at greater risk for food-borne illness. Most food-borne disease results from poor food handling.

To handle food safely:

- Keep food at safe temperatures. Bacteria grow quickly between 40° and 140°F (4° and 60°C). Keep cold foods below 40° (4°C) (refrigerated) and hot food above 140°F (60°C).
- Don't defrost food at room temperature. Defrost food in the refrigerator or place in cold water in a watertight bag change the water every 30 minutes. You can thaw food in the microwave if you cook it immediately after thawing.
- When thawing in the refrigerator, make sure meat juices do not contaminate other foods.
- Cook all meat and poultry thoroughly, to an internal temperature of 165°-180°F (73°-82°C). Use an instant-read thermometer. If you don’t have one, visually assess doneness; ground meat should not be pink and poultry juices should run clear, without any trace of red, pink or cloudiness.
- Use up leftovers within two days.
- Keep foods covered, wrapped or in airtight containers.
- Don’t eat raw eggs or foods containing them.
- Eat thoroughly cooked eggs only.
- Don’t leave hot or cold food out at room temperature for more than two hours.
- You may eat meats that have been appropriately refrigerated or frozen as long as they’re cooked adequately.
- Avoid cross-contamination of cooked meats with raw foods.
- Divide large amounts of food into small, shallow (2-4 inch) containers for quick cooling in the refrigerator. Cool hot foods uncovered in the refrigerator; cover once cooled.
- Refrigerate only as much as can be eaten in two to three days. Freeze the rest.
- Wash fruits and vegetables thoroughly under running water before peeling and cutting. Avoid any produce with mold.
- Wash tops of canned foods before opening and clean the can opener after each use.
- Never taste food that looks or smells strange. When in doubt, throw it out.

Kitchen Hygiene

- Thoroughly clean cutting utensils, countertops and cutting boards before and after each use.
- Use paper towels (not sponges) to clean up juices from raw meat or poultry. Sanitize the area.
- Use separate cutting boards (plastic, glass or wooden) for cooked or ready-to-eat foods and raw foods. Wash cutting boards after each use in hot soapy water or in the dishwasher.
- After cleaning, sanitize equipment and surfaces, including cutting boards, with dilute bleach solution: 1-3 teaspoons bleach in 1 quart or liter of water.
- Keep appliances, kitchen counters and other preparation areas free of food residue and particles.
- Clean cutting utensils, countertops and cutting boards before and after each use.
- Dry hands with paper towels. Replace dishcloths and dish towels daily and sponges at least once a week.
- Use an instant-read thermometer. If you don’t have one, visually assess doneness; ground meat should not be pink and poultry juices should run clear, without any trace of red, pink or cloudiness.
- Use up leftovers within two days.
- Keep foods covered, wrapped or in airtight containers.
- Don’t eat raw eggs or foods containing them.
- Eat thoroughly cooked eggs only.
- Don’t leave hot or cold food out at room temperature for more than two hours.
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- Refrigerate only as much as can be eaten in two to three days. Freeze the rest.
- Wash fruits and vegetables thoroughly under running water before peeling and cutting. Avoid any produce with mold.
- Wash tops of canned foods before opening and clean the can opener after each use.
- Never taste food that looks or smells strange. When in doubt, throw it out.

Microwave Cooking

Microwave cooking can leave cold spots in food where bacteria can survive. If your microwave doesn’t have a turntable, rotate the dish a quarter turn once or twice during cooking. When heating leftovers, partially cover food with a microwave-safe lid for thorough heating; stir several times during heating.

Refrigerator and Freezer

- Keep shelves and doors sanitized.
- Maintain refrigerator temperature at less than 40°F (4°C) and freezer temperature less than 0°F (-17°C).
- Discard foods that have exceeded their expiration dates. Discard all prepared foods after three days (72 hours).
- Discard any package or container that has moldy foods.

Cupboards and Pantry

- Discard without tasting any food that comes in a can or jar that is bulging, leaking, cracked or damaged at the seam.
- Discard any items past the expiration date.
- Make sure food storage areas remain reasonably clean, without insect or rodent contamination.
- If you have home-canned foods, review the processing procedure and conditions to be sure it was appropriate for the type of food. If you’re unsure about any item, discard it.

Grocery Shopping

- Avoid unlicensed sidewalk or roadside vendors.
- Check sell-by and use-by dates. Check packaging dates on fresh meats, poultry and seafood.
- Select undamaged produce. Check for unusual odor and mold or insect contamination.

Wash fruits and vegetables thoroughly under running water before peeling and cutting.

Vitamins and Herbal Remedies

If you’re currently taking any vitamins, nutritional supplements, cannabinoids/tetrahydrocannabinol (CBD/tet), or any herbal remedies, discuss them with your transplant doctor.

Do not take these herbal or botanical medications under any circumstances:

- Alfalfa
- Borage
- Chaparral
- Coltsfoot
- Comfrey
- Ephedra or Ma Huang
- Groundsel or Life Root
- Yohimbine
- Heliotrope
- L-tryptophan
- Lobelia
- Mate tea
- Sassafras
- Yohimbine

Dining Out

- Be sure that any food you eat is adequately cooked and freshly prepared. Avoid raw, uncooked and undercooked foods (sushi, steak tartar). All meat should be thoroughly cooked.
- Don’t order anything containing fresh fruits or vegetables. For example, avoid salad bars. Order hamburgers with a bun, meat and cheese only.
- Use condiments like ketchup and mayonnaise from individually wrapped packages.
- Pizza and fried foods are fine.
Long-Term Follow Up

Clinic Visits
You’ll need to stay close to your transplant center for at least 100 days after discharge. Your doctor will decide how long you should remain in the area.

You’ll return to see your transplant doctor at days 30, 100 and 180, and then annually. Please keep your scheduled clinic appointments. If for any reason you’re unable to attend, call the transplant doctor’s office.

Immunization Schedule
Your immune system is your body’s defense against infection and disease. After having a stem cell transplant (SCT), you will probably have a weakened immune system. This means you’re at a higher risk for infection and disease. Your team will give you your vaccine plan when you are ready.

You’ll receive the following immunizations only if you don’t have active, chronic GVHD (see page 17) or ongoing immunosuppressive treatment at time of vaccination.

- Covid
- Diphtheria, tetanus, pertussis (DPT)
- Haemophilus Influenzae Type B
- Hepatitis B
- Influenza
- Measles, mumps, rubella (MMR)
- Pneumococcal
- Pneumovax (or Prevnar if immunosuppressed and chronic GVHD)
- Poliovirus
- Shingles

Information and Education
- American Cancer Society: cancer.org
- American Society for Blood and Marrow Transplantation: asbmt.org
- Association of Cancer Online resources: acor.org
- Blood and Marrow Transplant Information Network: bmtinfonet.org
- Bone Marrow Foundation: bonemarrow.org
- Cancer Care: cancercare.org
- Cancer Net: cancernet.org
- International Myeloma Foundation: myeloma.org
- Leukemia and Lymphoma Society: leukemia-lymphoma.org
- National Bone Marrow Transplant Link: nbmmlink.org
- National Cancer Institute: cancer.gov
- Sutter Cancer Services: sutterhealth.org/cancer

Exposure to the live polio vaccine is dangerous. Avoid contact with children or adults who’ve received the live polio vaccine in the previous three months.

Online and In-Person Support

Everyone has different needs as they progress through the BMT process. You may find many programs and resources useful as you consider what you need. Your BMT team is available for questions and to help you secure resources. Your social worker, financial coordinator, NMDP coordinator and transplant coordinator are reliable sources of guidance and support.

My Health Online: mho.sutter.org
- Access your Sutter Health electronic health record.
- Request medical appointments.
- Message your care team.
- View lab test results.
- Request prescription renewals.
- Access health information.

Information and Education
- American Cancer Society: cancer.org
- American Society for Blood and Marrow Transplantation: asbmt.org
- Association of Cancer Online resources: acor.org
- Blood and Marrow Transplant Information Network: bmtinfonet.org
- Bone Marrow Foundation: bonemarrow.org
- Cancer Care: cancercare.org
- Cancer Net: cancernet.org
- International Myeloma Foundation: myeloma.org
- Leukemia and Lymphoma Society: leukemia-lymphoma.org
- National Bone Marrow Transplant Link: nbmmlink.org
- National Cancer Institute: cancer.gov
- Sutter Cancer Services: sutterhealth.org/cancer

Peer Support
- Sutter BMT Patient Contact Coordination. For one-on-one contact with a previous Sutter BMT patient or caregiver, please call your BMT social worker or transplant coordinator to match you with an appropriate candidate. This might include contacting someone by phone or in person depending on your comfort level.
- Sutter Support Groups. Sutter Cancer Services offers support groups for cancer patients, family and caregivers, plus bereavement and art therapy for adults and children. For a complete list of groups, locations and times, visit sutterhealth.org/cancer and click on “Special Services and Support.”
- Counseling Services. Sutter Cancer Services provides free counseling with a licensed professional who specializes in cancer patients and their families. Ask your social work team for more information.
- On-Call Peer Support. Get free, one-on-one support from trained peer support volunteers who are BMT transplant survivors, caregivers and donors. To request this, call 800-LINK-BMT (800-546-5268) and ask for peer support.
- Supportline. Receive free online or phone support from a BMT patient or caregiver through the Bone Marrow Foundation. Call 800-365-1336 or fill out a request at bonemarrow.org/supportline.

ACOR Online Communication Forums. The Association of Cancer Online Resources offers multiple online communication forums, including lists specific to BMT and caregivers. After creating a login, you’ll have access to monthly newsletters and a listserve for posting and communication. Go to acor.org to start.

Create Your Own Website or Blog
- CaringBridge.org
- LiveJournal.com
- MyLifeline.org
Financial Resources

Several local and national organizations offer financial assistance for eligible patients with a cancer diagnosis. In addition, many pharmaceutical companies have programs for those who cannot afford medications. Income replacement options for you and your caregiver(s) include State Disability Insurance, Social Security Disability or Paid Family Leave. Ask your employer for information about your benefits and eligibility for Family and Medical Leave Act (FMLA).

Many programs require documentation of your diagnosis or treatment plans as well as proof of your financial need. Your BMT social worker or financial coordinator can offer information about these programs, including private fundraising, and assist with applications.

- American Cancer Society: cancer.org, 800-277-2345
- The Bone Marrow and Cancer Foundation: bonemarrow.org
- California Employment Development Department (EDD): ca.edd.gov
- CancerCare: cancercare.org, 800-813-4673
- Leukemia and Lymphoma Society: lls.org
- Social Security Administration (SSA): ssa.gov, 800-722-1213

Applications for Financial Assistance

Please tell your transplant coordinator if you need any of these forms. Some require a healthcare provider’s signature.

- American Cancer Society Patient Referral Form
- The Bone Marrow Foundation Patient Aid Program Application
- DMV Application for Disabled Person Placard or License Plates
- Your local utilities may offer financial assistance. Contact them directly.

Transportation

Ask your Coordinator for local resources to assist you with any transportation needs.

Local Housing

If you can’t remain at home through the transplant process, please discuss your needs with your social worker. Your BMT team can help if you need proof of hospitalization or treatment.

The American Cancer Society collaborates with hotels around the county to offer temporary, free or discounted rates for cancer patients who need to travel for treatment. Call 800-227-2345 for additional information.

The Cancer Support Community collaborates with Airbnb to offer temporary, free housing for income-eligible cancer patients who need to travel more than 50 miles from home for treatment. Discuss this option with your social worker, call 877-793-0498 or email helpline@cancersupportcommunity.org for more information.

Sutter has negotiated discount rates for our patients at several local hotels. Please go to suttermedicalcenter.org/housing for a current list of hotels and rates.

Visit Joe’s House, a Lodging Guide for Cancer Patients, at joeshouse.org for links to local hotels that offer discounts to cancer patients.

For rental apartments, RV accommodations and other options, talk to your BMT social worker.

Recommended Books

The Anatomy of Hope: How People Prevail in the Face of Illness
Jerome Groopman, M.D.

Becky and the Worry Cup: A Children’s Book About a Parent’s Cancer
Wendy S. Harpham, M.D.

Climb Back from Cancer: A Survivor and Caregiver’s Inspirational Journey
Cecilia Hobson and Alan Hobson

Everyone’s Guide to Cancer Supportive Care: A Comprehensive Handbook for Patients and Their Families
Ernest Rosenbaum, M.D., and Isadora Rosenbaum

The Human Side of Cancer: Living with Hope, Coping with Uncertainty
Jimmie C. Holland, M.D., and Sheldon Lewis

My (So-Called) Normal Life
Erin Zammett

Picking Up the Pieces: Moving Forward After Surviving Cancer
Sherri Magee, Ph.D., and Kathy Scalzo, M.S., O.D.

When a Parent Has Cancer: A Guide to Caring for Your Children
Wendy S. Harpham, M.D.