Taking care of yourself

Caregivers are at high risk of developing depression and other health problems.1 Caregiving is sometimes very demanding and it may be difficult to find time to consider your own wellbeing. If you are burnt out, exhausted or ill it will be much harder to provide support.

Below are some suggestions for ways to take care of yourself while caring for the person with bipolar disorder:

• **Become informed:** Become informed about bipolar disorder and ways to treat and manage it.

• **Give yourself time to adjust:** (see Box 1: Coming to terms with the person’s bipolar disorder). Caregiving situations differ and finding what works to deal with the person’s bipolar disorder can be a trial and error process.

• **Use helpful resources:** Make use of appropriate services and community organizations to support the person, yourself and the family (e.g. health, financial or respite services, and peer support groups). If it is hard to manage work and the demands of the illness, find out about possibilities for leave and flexible working hours.

• **Make advance plans to deal with the bipolar disorder:** Plan ways to deal with bipolar episodes or crises in advance (see bipolarcaregivers.org for making plans and agreements).

• **Use problem solving to sort out difficulties:** Using a problem solving approach to sort out difficulties can make them less overwhelming (see Box 2 for problem solving steps).

• **Use stress reduction strategies when the person is ill:** Patterns of illness can vary and there may be more caregiving demands when the person is experiencing a bipolar episode, if they have persistent symptoms or they relapse frequently (see Box 3 for reducing stress when the person is ill and the summary on ‘Dealing with bipolar crises’).

• **Maintain contact with friends and family:** Isolating yourself from social contact can contribute to depression. Having someone to talk to whom you can trust can make it easier to cope. Caregivers may also benefit from professional counseling.

• **Maintain healthy boundaries with the person and their illness:** As far as possible, try not to focus on the person’s bipolar disorder all the time. Maintaining your own interests, activities and relationships can help to keep things in perspective and prevent exhaustion and burn out. See Box 4 for more suggestions about maintaining healthy boundaries.

• **Keep healthy:** Despite the demands of caregiving try to maintain some healthy lifestyle habits (see Box 5) and recognize and respond to your own signs of stress (see Box 6) or depression (see Box 7). If you notice signs of stress or depression, don’t criticize yourself for these understandable reactions. Develop a plan for coping (e.g. delegate more of your duties and take time off). If you have symptoms of depression, contact your doctor for an assessment and to discuss treatment options. Don’t ignore depression.

• **Develop realistic expectations:** Sometimes caregivers have unrealistic expectations about what they ‘should’ do to help (see Box 8 for a few examples of unrealistic expectations and more realistic alternatives). If you feel out of your depth when trying to help the person, say something supportive that acknowledges the need for extra help. Having unrealistic expectations of what the person and their clinician can do to control bipolar disorder can also lead to frustration and disappointment. While there is a lot that they can to prevent relapse, bipolar episodes sometimes occur despite everyone’s efforts to prevent them.
• **Acknowledge your natural reactions:** Bipolar symptoms and their consequences can evoke a range of intense emotions (e.g. anger, sadness, guilt, shame, feeling rejected or like you want to withdraw from the person or to leave). Acknowledging these natural reactions and deciding how to deal with them can reduce stress. For example, if you are frequently angry, find a constructive release for your anger (e.g. go for a walk, play sport, paint, write in a journal, or talk things through with someone you trust). If something the person you care for has done has made you angry, wait until you have calmed down before discussing this behavior with them. If possible, the best time to discuss this is when the person is relatively well. For more about dealing with your natural emotions see bipolarcaregivers.org.

• **Maintain or rebuild your relationship with the person:** Relationships are sometimes affected by the person’s bipolar disorder and the caregiving situation. There might be times when the person is too ill to respond to you in the way you would like them to (emotionally, or where partners are concerned, sexually). See box 9 for information about maintaining relationships.

• **Recognize the positives:** Some caregivers find positive things about caregiving that make it a bit easier to cope with the person’s extreme bipolar moods. For example, they realize that through caregiving they have become more tolerant empathic and gained more confidence, or they notice positive things about the person such as their creativity or courage in trying to manage their bipolar disorder.

• **Acknowledge the support you provide:** The tremendous support caregivers provide sometimes goes unrecognized. Acknowledge the effort you make and the care you provide.

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**BOX 1** Coming to terms with the person’s bipolar disorder

Caregivers often experience a range of understandable reactions when trying to come to terms with the person’s illness (e.g. ranging from initial shock, disbelief and emotional turmoil, to gradual understanding, acceptance, and hope that there are ways to deal with bipolar disorder and live well). This grief often subsides. However, emotional turmoil may return, for example when the person relapses.

If you experience emotions linked to this natural grief process:
- Allow yourself to grieve.
- Reassure yourself that while the changes and losses connected to the illness are real. Despite the illness, people with bipolar disorder and caregivers often find new meaning and ways to enjoy life.
- You might find it helpful to make contact with others in similar situations who can relate to what you are going through.
- Consider setting small goals to do things that you enjoy.

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**BOX 2** A problem solving approach

There are 4 problem solving steps when sorting out difficulties:

**Step 1: Clearly define the problem**
Without blaming yourself or others, consider how the problem developed, when it occurs and why it is a problem.

**Step 2: Decide what solution or solutions to try**
First make a list of the possible solutions. It doesn’t matter how unrealistic the solutions are at this stage. Next, work out how practical or realistic each solution is in your situation. What are the possible risks and negative consequences that may occur if you choose this solution? Are there ways to prevent or deal with these consequences if they occur? What are the possible benefits if you choose this solution? Finally select a solution or a few solutions to try.

**Step 3: Develop a plan of action and follow the plan**
Decide what you need to do first to put the solution(s) into practice and work out a step-by-step plan. Put your plan into action.

**Step 4: Review how the solution worked**
Acknowledge the effort you have put into trying to manage the problem. Give yourself credit if it worked. However, some problems are more difficult to solve than others. If the solution was not helpful or there are still parts of the problem that need to be solved, return to Step 2 to try other solutions. Difficult situations can take time to change or may not even be able to be changed. If this is the case, look for ways to make things a little easier and enjoyable for yourself, despite the situation.

Finding hope - Sarah K Reece
BOX 4  Maintaining healthy boundaries

- Acknowledge that you have needs too.
- Keep in mind that although you can help, the person needs to find ways to deal with their own illness.
- Set realistic limits on what you can do to help by considering the severity of the person's symptoms, your own wellbeing and other commitments and who else can help (e.g. family, friends, clinicians and community organizations).
- Learn to say “no” to demands that are unreasonable or unmanageable (both caregiving and other demands).
- Devote some time to outside interests and goals; doing something that you enjoy.
- Encourage the person's efforts to maintain their own identity and interests.
- Use the time when the person is well to focus on things that are important to you, besides the illness.
- Where to set limits with risky, dangerous or inappropriate illness behavior is a personal choice (see ‘Dealing with risky or inappropriate manic or hypomanic behavior’ on bipolarcaregivers.org).

BOX 5  Maintaining a healthy lifestyle

To improve your lifestyle, rather than trying to do everything at once, set a goal to include something from the following list and gradually change your lifestyle:

- A little regular exercise
- Eating healthy balanced meals
- A bit of regular time off to relax or pursue a hobby
- Time off to keep in contact with selective friends, family or a peer support group
- Regular sleep
- Treatment if you have a health condition.
- Reduce ways of coping that are destructive to your health (e.g. drinking too much alcohol, smoking, overeating when stressed, neglecting your hygiene to save time, isolating yourself from others as they may not understand your situation).

BOX 6  Signs of stress

- Being more irritable, short-tempered tense or anxious than usual
- Overeating or loss of appetite
- Difficulty falling asleep
- Having frequent minor illnesses or aches and pains or feeling run down
- Using alcohol or other drugs to cope.
BOX 8 Examples of unrealistic expectations of yourself as a caregiver and realistic alternatives

<table>
<thead>
<tr>
<th>Unrealistic expectation</th>
<th>Realistic expectation</th>
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<tbody>
<tr>
<td>“I should cure the person’s bipolar disorder” or “I should be able to fix everything”.</td>
<td>Bipolar disorder is a complex illness that needs ongoing management, rather than something that can be fixed forever. While you can be supportive, it is the person’s illness and their responsibility to manage it. A more realistic alternative might be: “There are things I can do to support the person to deal as best as they can with their bipolar disorder.”</td>
</tr>
<tr>
<td>“I should always be perfectly supportive of the person and never feel stressed or angry as they are ill.”</td>
<td>A more realistic expectation might be: “Although there are things I can do to be supportive, no one can be perfectly supportive all the time. It is understandable that I feel stressed or angry at times.”</td>
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BOX 9 Maintaining your relationship with the person

- When the person is ill, try to separate the bipolar disorder talk and behavior from the person. Try to detach from the situation. Don’t take the sometimes hurtful things the person says when they are ill personally. However, set limits with verbal abuse. For example, if the person is verbally abusive, consider saying something like “I understand you’re upset but I’m not going to tolerate being spoken to in this way” and walk away.
- When the person is less ill do things together that you both enjoy.
- Whenever possible, encourage ‘give and take’ in your relationship.
- Don’t always make the person’s bipolar disorder the focus of the conversation between you. When possible, relate to the person as the friend or family member they are.
- Suggest to the person that you make plans together about ways to deal with the bipolar disorder to reduce its impact on both of you.
- Problem solve around difficulties but also focus on sharing positive things (e.g. personal achievements or pleasurable experiences).
- There are ways of communicating with each other about grievances that are constructive and are less likely to come across as hostile or critical. For example, listening carefully to the person’s point of view, making positive requests for change and calmly expressing your feelings about their behavior (see good communication skills on bipolarcaregivers.org).
- If there is a lot of conflict between you, consider consulting a relationship counselor with experience in working with bipolar disorder.
- If the person is physically aggressive: Physical aggression rarely occurs as part of bipolar disorder. It is more often connected with drug or alcohol problems, personality disorders and occasionally with psychosis. However, occasionally if a person who is manic or in a mixed episode is very angry, they may act out on their emotions. Never compromise your own or others’ safety due to concerns about hurting the person’s feelings, as later the person might feel very relieved that they were prevented from hurting their loved ones. Make sure you are safe first and contact the emergency services. See box 10 for more about protecting yourself if the person you care for has previously become aggressive.

BOX 10 Ways to protect yourself if the person has become aggressive before

- Learn to recognize the warning signs of impending aggression.
- Take even casual threats of violence seriously.
- Work out in advance how to ensure your safety and that of others (e.g. have locks on rooms, leave the house and get help when warning signs of aggression appear).
- Remove objects that could be used as weapons if the person is likely to become aggressive.
- Use the services available to assist you to put safeguards in place to prevent this abuse.
- It is common to feel very traumatized by physical abuse and professional counseling can be helpful.

References