Role, Health and Well-being

An inspirational guide for care partners of people living with dementia

SERIES 3
Introduction

This By Us For Us guide is the second in a series of helpful resources written BY and FOR care partners of people living with dementia. A care partner can be anyone including a spouse, child, sibling, grandchild, extended family member, neighbour, legally appointed representative, member of a community with whom a person living with dementia may be associated (e.g., spiritual group, ethno-cultural group), and so on.

This new care partner series is based on the original By Us For Us series written BY and FOR people living with dementia. This series is a collaboration of care partners, many who were interviewed and several more involved in the research, development, and editing process.

This guide outlines some roles that care partners may assume and the potential impact on both the care partner and the person living with dementia. Tips and strategies are offered to assist you with understanding, supporting, and maintaining your health and well-being. Care partners share with you the positive insights they have gained along their journey.

In this guide you can find valuable information about:

1) The many roles that care partners take on.
2) The difficulties and challenges experienced by care partners.
3) Effective strategies from care partners for managing and coping with difficult and challenging experiences.
4) Stories of positive experiences along the journey.

– Emma, Charlene, Jack, Karen, Lisa, Kimberly, and Jessica

The By Us For Us Care Partner Committee
For those living with a form of dementia, loss and changes in abilities are expected. When dementia begins to affect specific abilities of a person, the care partners will most likely need to take up these responsibilities. As such, the role of the care partner changes accordingly. Being a care partner does not just mean providing hands-on care to people living with dementia. Being a care partner means taking on new and multiple roles. These roles continually change and can impact the health and well-being of the care partner. When some care partners take on new roles, they find it challenging and overwhelming, while others talk about it as being rewarding and providing opportunities for personal growth.

Some of the new and unexpected roles that a care partner may take on can be easier or more natural to fill than others. This may depend on:

- the relationship with the person living with dementia,
- past experiences,
- the level of support needed by the person living with dementia,
- progression of dementia, or
- available resources.

For example, you may find yourself cooking or managing finances for the first time.
You may find yourself helping with personal care, assisting the person living with dementia with tasks such as dressing, bathing, and grooming. These tasks are often referred to as “Activities of Daily Living” (ADLs). Your role may expand to being a coordinator, who plans appointments and meetings related to healthcare, social, and leisure activities. You could find yourself being an educator; teaching family, healthcare professionals, the public, and others about dementia. Individuals in the support circle, including people living with dementia, may also need you to provide emotional reassurance.

You may find yourself being an advocate. This role could consist of explaining the needs of the person living with dementia when they are unable to communicate for themselves. It may also involve finding ways to help the person living with dementia obtain the services and programs needed in order to maintain their well-being, comfort and dignity. If you are appointed lead decision maker (also known as ‘Substitute Decision Maker’), you may become legally responsible for making decisions related to finances and/or healthcare.

“My role has evolved from being a secondary caregiver, primarily giving advice when my father was alive and able to care for my mother and in the early stages of dementia when my mother was quite independent, to being the primary caregiver who has to make all financial decisions, and many caregiving decisions on behalf of my mother.”

(Care partner)
Wearing of many “hats” continued

This may involve decisions regarding real estate or medical treatment. Some care partners take a more pro-active role by publicly sharing their personal stories on how they live well in the face of dementia to challenge the stigma.

These roles contribute to an increase in your responsibilities and workload. Regardless of the roles you take on as a care partner, **the most important thing to remember is not to lose yourself in this journey.** You need to maintain your own identity, and continue to look after your own health and well-being in order to provide the best care and support possible to your family member or friend living with dementia.

**Use the space below to indicate how your role has evolved and what ‘hats’ you wear.**

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Impact of changing roles

Changing roles may have an impact in many areas for care partners. A caring role may lead to increased stress, illness, emotional and/or physical fatigue, and even decision fatigue. Decision fatigue can create feelings of inadequacy in making decisions, a desire to make decisions too quickly just to get it over with, or a sense of exhaustion. The following are several more areas that can be impacted.

Nature of relationships

Changes in your roles can sometimes affect the nature of the relationship between you and the person living with dementia you are supporting. For example, an adult child may feel overwhelmed or uncomfortable by having to help a parent with dressing or bathing.

Spouses may experience a sense of loss of companionship as their partner’s dementia progresses and roles begin to change. Sharing life experiences with the person living with dementia may become more difficult as dementia progresses. Maybe you used to plan and share holidays together and are no longer able to do so. This can be accompanied by frustration and a feeling of sadness or grief about this loss, making the situation hard to accept. As a result, it can be more difficult for you, as a care partner, to provide the care the person living with dementia requires.
Financial impact

There may be financial changes such as the need for additional paid support, medications, tests, and long-term care fees. For families, income may also be reduced due to early retirement, loss of employment or modified work.

Lifestyle changes

You may experience a loss of freedom. You could be faced with your own physical and/or mental challenges. Some of your friends may distance themselves if they are uncomfortable with dementia and its effects.

Feelings, mood and emotions

Travelling the journey with someone living with dementia can cause a large range of emotions including sadness, anxiety, fear, and frustration. For the care partner, these can be a natural consequence of changing roles.

“My role as a partner in care has gone from just being around for company for my partner, to having to check to see what he is doing fairly frequently… I have to drive him places, help him with the ATMs… and solve problems for him.”

(Care partner)
Feelings of frustration and anger about changes in the capabilities of the person living with dementia, or the increased need for support, can also affect the health of the care partner. You may not know which steps to take, or you may lack confidence about managing unfamiliar skills. Friends and family may offer a variety of solutions for the challenges you are facing. However, they may lack understanding or provide unrealistic solutions; you could be left feeling isolated or trapped.

A person living with dementia becomes increasingly aware of your tone, mood, and body language and may have challenges interpreting your spoken words. Your mood and body language can significantly affect a person living with dementia. If you are positive, they are likely to reflect that and be positive themselves. If you are negative, they are likely to reflect that and be negative themselves. This is known as the “plasticity” of dementia. It can be difficult to regulate one’s own attitude, especially when the person living with dementia is a spouse.

Sometimes you may feel frustrated that the person living with dementia does not understand what you’re saying, so you automatically raise your voice. However, this is a challenging situation as the problem may actually be comprehension and not hearing.

“When things turn around to more bad days, things change and it becomes very worrisome and you start to fatigue. That’s when you feel the most pressure and alone in the world.”

(Care partner)
A loss of a capability, even a small one, causes grieving. Denying a loss just makes the feelings surface somewhere else in a less manageable way. Your friends may not have much experience with grieving the kinds of gradual losses that occur when a person is living with dementia. They may have more experience with grieving a sudden loss, such as a death, for example. This may make it more difficult for them to grasp your feelings of loss and isolation.

Compassion fatigue is a term used to describe a form of burnout that professionals, volunteers and care partners can develop while helping others in need. Helping professionals are individuals such as nurses, doctors, police officers, and social workers. Care partners, volunteers, and professionals may find that over time they feel less sympathetic or patient with the stories they hear from the people they work with, and become less compassionate towards their friends and family.

Care partners may extend themselves beyond their own capabilities and may not be able to recognize their own limitations, making them less effective in the quality of care they provide. With compassion fatigue, care partners, helpers, and volunteers start feeling very emotionally and physically exhausted. Those with compassion fatigue can lose their enjoyment of the task and develop a more negative view of their world.
It can be challenging to find a balance between your own life and the responsibilities associated with your new roles. One of the weaknesses of our amazingly capable human body is our poor ability to detect very gradual changes in ourselves. As the roles for the care partner shift and responsibilities increase, you may continue to think that you are adapting well to handling the load. You may find yourself defensive about how you are doing. The phrases, “I’m OK” or “I’m fine and handling things well” may be your answer. Others observing you may be better able to see your stress level, your exhaustion, or the effects of your feelings of isolation.

Use the space below to indicate how your role has evolved and what ‘hats’ you wear.

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It is important to understand that this journey may be life altering (i.e., physically, emotionally, intellectually, spiritually, etc.). Once you’re in a place of acceptance, it becomes easier to continue through your journey.

Do not wait until a crisis happens to seek support for yourself! It will help if you develop a good understanding of dementia and the journey that both you and the person living with dementia will share. Educating yourself and planning ahead can help you not only cope better, but will allow you to maintain a sense of well-being despite all the changes that you experience.

The following are some general tips and strategies that can help you cope with the changing roles and minimize the impacts to your health and overall well-being.

Understanding our health and well-being and the journey ahead

Educate yourself and others

Become more knowledgeable about the illness. Talk to your doctor and/or your local Alzheimer Society. Information found on the internet and through support groups can also be beneficial. However, be careful about the information you find online; it may not always be from a reliable source.
Do not take changes personally

There are many factors that create changes in the person living with dementia including physical well-being, social changes and their environment. Their actions and words change because of the dementia and are not meant to hurt you, but rather as a result of the dementia process. It is important to put the blame on the illness NOT the person.

Know your “breaking point”

“Breaking point” is the moment when stress gets so overwhelming that you can no longer cope or manage a situation. Your “breaking point” may sneak up on you because these changes can happen very gradually. Listen to the advice of your support network objectively. Use professional/counselling support services before you get to your breaking point. Remember, the quality of care that we can deliver is directly related to our own health.

Do advanced care planning

It is important to think about the future as early as possible in the progression of dementia. This will ensure that the wishes of the person living with dementia will be taken into account later. The more familiar you are with the wishes of the person living with dementia, the easier it is to have confidence in making decisions on their behalf in the future. This could also reduce the risk of conflict with family members or others supporting the person living with dementia.
General tips and strategies continued

✔ Draw on available informal supports – engage family, friends, neighbours, or anyone who can provide positive support to build your informal network.

✔ Search out available formal supports – there may be support services available in your community. Find a format that works for you (i.e., respite, support groups, etc.). Learning about them helps you plan and build your support network.

✔ Make financial plans – become familiar with financial options that can help manage future challenges, such as reduced employment income and costs for respite and long-term care.

✔ Discuss health care – talk openly about health care and support including preference for living accommodations (i.e., placement in a long-term care home, waitlists, etc.), quality of life choices, and end of life care (e.g., “Do Not Resuscitate (DNR)” orders, continuation of medication for other illnesses).

“Struggles are there on your journey and by getting educated and reading, talking about it seems to help.”

(Care partner)
General tips and strategies continued

Supporting health and well-being through positive behaviours

Acknowledge and accept your feelings and emotions

Don’t blame yourself for the changes in your mood. Even for small losses, allow yourself to go through the grieving process. By doing so, care partners often find themselves in a better place of acceptance and even develop a sense of peace and hope.

For more information on grief and loss, refer to the By Us For Us guide: Living and Transforming with Loss and Grief.

Take care of yourself

Make sure to get plenty of rest, healthy nourishment, and exercise whenever possible. Taking good care of yourself is important in helping you to provide effective support for a person living with dementia. It also reduces your risk of compassion fatigue.
Accept help from others

When family or friends offer to help, it may be very valuable to accept their assistance. Sharing some of your roles with others allows them to become familiar with the situation. It also allows the person living with dementia to become comfortable accepting assistance from others. Drawing on people whose opinions you respect, those with more experience, or those you know well will help you to avoid decision fatigue.

Take time to get away and nourish your soul

Try using adult day programs, respite services, and your support networks so you can maintain life outside the caring role. Avoid self-isolation by staying personally active (e.g., support groups, personal care, theatre, travel, shopping). This is not something that you should feel guilty or selfish about. You are doing it for the person living with dementia so you can be a better care partner.

Adopt humour and a positive outlook in your daily life

Take time to laugh. It is healthy to laugh at surprising situations. Taking on a positive attitude is a good thing. A cheerful voice, a smile, or some uplifting words can make a big difference. Appreciate the emotional connection you may have with the person living with dementia. Focus on living in the moment and the quality time you have together.
Maintaining health and well-being in a changing reality

Recognize and support abilities

It is important to recognize and support the changing capabilities and strengths of the person living with dementia. This will help foster a better quality of life through the journey. It is also important to try to understand their reality and let them live in it. Don’t argue or try to control their perspective.

Be as flexible as you can

Each day may differ. Things may not go as planned… that’s to be expected. Live in the moment and remember some things may be beyond your control.

“You] ...need to be realistic. We cannot prevent the disease progression so how can we minimize impact – take care of yourself (physically, mentally, emotionally, and spiritually)!”

(Care partner)
General tips and strategies continued

Use time saving strategies to balance hectic schedules

As the dementia journey changes, your schedule may become busier. Many care partners have found organizers help them stay on track. Figure out what strategies work for you. It can be helpful to prioritize, make lists, or delegate tasks to others.

Listen to respected opinions about your well-being

If trusted family, friends, and neighbours are expressing concerns about your well-being, remember that they can often detect gradual changes in yourself better than you can.

Recognize that you are not alone

Many have experienced the same feelings and emotions you may be facing. Draw on their knowledge and expertise throughout the journey.

“I have learned to be very patient and go with the flow. I try to make my partner’s days the best they can be by altering my behaviour to fit with the ‘new’ person I am living with.”

(Care partner)
In the midst of the journey, positive experiences can be difficult to recognise. However, many care partners talk about the rewards they have gained. In fact, focusing on the positive is a helpful way of coping. Some recognise the new skills, insights, and experiences that are learned along the way. Many of these skills can be transferred easily to other new life situations.

The following are some of the positive experiences or insights that care partners have gained along the journey:

- Development of new skills, expertise, confidence, and a realisation of one’s own limitations and when to ask for help.
- Continued appreciation of the person living with dementia and their strengths.
- Reflection on goals, priorities, and values in life.
- Moments of thankfulness and blessing.
- Enhanced ability to appreciate life and its many pleasures (e.g., sunrise/sunset).

“I have [now] become proficient at planning and preparing meals, doing the laundry, ironing, grocery shopping, shampooing, and setting hair, nails polishing, and learning [about] the [challenges] of sizing ladies bras and underwear.”

(Care partner)
Being present and sharing each other’s company (i.e., warm smile, embrace, or holding hands).

Strengthened faith or spirituality.

Improved quality of relationships with other care partners, family, and friends (e.g., stronger network or relationship repair).

Stronger emotional connection with the person living with dementia (e.g., by listening to their stories, feelings, and experiences).

A sense of accomplishment, resourcefulness, and pride in doing the best that you can.

Greater acceptance of and sensitivity to others.

Pleasure in being able to reciprocate love and care.

Care partners can learn to appreciate all that people with dementia may have to offer and the joy in day-to-day life. This may involve relying on professional help, a change in mindset, a reliance on spiritual supports, or taking time to nourish the soul.

“... [I] am learning to appreciate the small things, the times that we get along nicely, and to live [in] the present, and not to worry about the future, which I can’t control (for the most part).”

(Care partner)
“From my care role, I have learned to appreciate the gift of my partner and the gift of all those who care and share our journey.”
(Care partner)
“Actually, I praise God more and more for the wonderful wife He has given me and for all the opportunity that this disease had given me to love her more.”

(Care partner)
When Brenda Hounam was diagnosed with Alzheimer’s disease at the age of 53, she realized that very little information was available for people living with dementia. The dominant view was that people living with dementia could no longer learn and be involved in their own care. Brenda knew differently. In 2003, she was inspired to address this gap - to develop a series of resources specifically designed by and for people living with dementia. In 2004, she approached two of her peers with her idea and was surprised by the enthusiasm and support they offered. Through Brenda’s contacts at the Alzheimer Society of Brant, the Alzheimer Society of St. Thomas, and the Murray Alzheimer Research and Education Program (MAREP), Brenda connected with various people living with dementia from all around Ontario to work on what came to be called, the By Us For Us (BUFU) guides. These guides provide tips and strategies for managing daily challenges and enhancing well-being for themselves and others living with dementia. Since the publication of the first guide, “Memory Workout”, the following guides focused on issues raised by people living with dementia. The project has grown to include three series researched and developed in partnership with people living with dementia and care partners. The philosophy of the By Us For Us guides remains committed, placing importance first on the experiences, suggestions, and tips from those directly affected BY dementia, FOR those directly affected. Brenda has been a true inspiration and role model to her peers. Without her determination and perseverance, this project would not exist or have the international recognition that it does.
Resources

Alzheimer Society of Canada
Toll-free telephone: 1-800-616-8816
Email: info@alzheimer.ca
Website: www.alzheimer.ca

World Health Organization
Website: www.who.int

We also suggest that you check for other local, national and international resources.

Endorsements

When a spouse, family member or friend is affected by memory loss and dementia, many of us become partners in care. This role can be both deeply rewarding and very taxing. Over time, managing competing demands and the emergence of difficult symptoms in the person with dementia can lead to compassion fatigue in the partner in care. This is a normal consequence of being in the role of a partner is care. The sooner we can detect that we are experiencing compassion fatigue, the better we can address it before it takes a toll on us and the person with dementia. Congratulations to BUFU for this compassionate and supportive resource guide that provides partners in care with concrete tools to navigate the challenging journey they face in caring for a person living with dementia.

Françoise Mathieu, M.Ed. CCC.
Author of “The Compassion Fatigue Workbook” published by Routledge in 2012
At the age of 26 I became a secondary partner in care for my father who had been diagnosed with early-onset Alzheimer’s. During this time, I found that information from individuals who have lived a similar experience to be the most useful. Watching my mother as the primary partner in care, I could see when she needed time to care for herself, but not all care partners have that kind of support. The BUFU guide reminds me how we dealt with so many different situations and I wish we had something like this during our journey. It is an excellent resource for persons living with dementia and their families.

Chris Wynn
Care Partners, Documentary Filmmaker

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Front cover concept

The cover art for this guide was based on the idea that the care partner role can bring life-altering changes in response to an environment that is sometimes fluid but often dynamic. We may need to reach out for others and stretch ourselves beyond our current knowledge or levels of comfort. Maintaining the care partner’s physical, psychological, spiritual or social wellness is an essential factor for providing the strength to face new challenges – renewing us with the energy to embrace the days ahead.
About the By Us For Us Guides

The By Us For Us Guides (BUFU guides) are designed to provide people living with dementia/care partners the necessary tools to enhance their well-being and manage daily challenges. What makes these guides unique is that they are created BY people living with dementia/care partners, FOR people living with dementia/care partners. The guides include three unique series:

People Living with Dementia: Series 1
This series is written by people living with dementia and covers their experiences. It includes topics such as tips and strategies for managing daily challenges, memory exercises, and enhancing communication and wellness.

Partnership: Series 2
This series is written by both people living with dementia and care partners and covers the partnership experience. It includes topics such as safety, food and mealtimes, social connections, and coping with loss and grief.

Care Partner: Series 3
This series is written by care partners and covers their experiences of supporting a person living with dementia. This series includes topics such as before diagnosis, young care partners, roles, and health and wellbeing.

To view other guides or order print copies, please visit the Schlegel-UW Research Institute for Aging website: www.the-ria.ca

We welcome your input…
If you are living with dementia or are a care partner and would like to comment on this guide, suggest topics for future guides, or participate in future guides, please contact Murray Alzheimer Research and Education Program (MAREP).
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