Time for Living and Caring
Making Respite Services Work for You!

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Fifth Edition
Faculty at the University of Utah Gerontology Interdisciplinary Program began conducting research related to caregiving in 1984. Faculty at California State University, San Bernardino joined in this effort in 2009 when Dr. Lund relocated to California. Over these years, we have learned that many family caregivers are doing an excellent job of doing everything they can to help their loved ones. However, providing care can be difficult and challenging especially when these efforts can last for years. We also know that many caregivers are trying to do more than their abilities, circumstances and resources allow. Unfortunately, many of these heroic caregivers are risking their own personal well being and sometimes their family life in the process. Many caregivers are not fully aware of the short and longer term consequences of “doing it all alone” because they are so overwhelmed with managing their daily tasks and responsibilities.

We hope that you are not one of these family caregivers, but you may need to pause long enough to read this brochure and find out if you are. The purposes of this brochure are to:

1. Help family caregivers assess their abilities and circumstances
2. Help caregivers to become aware of sacrifices that they are making
3. Encourage family caregivers to seek the help of respite services
4. Provide suggestions to make respite most effective

We are convinced that when caregivers use respite services they are better able to preserve quality in their own daily lives and enhance overall family life. An additional benefit is that the loved ones in need of care usually get better quality care from their caregivers because the caregivers are more likely to have some balance in their lives and better health.
Avoiding Potential Problems

Before we describe some of the various forms of respite services, we would like to summarize some of the main findings that research has revealed about caregivers and what often happens when they do not get sufficient help with their caregiving tasks. You need to be aware of these possibilities so that you know what is happening in your own life. We want to help you become better informed so that you can make better decisions about your caregiving situation.

Although not all caregivers experience overwhelming stress, research has documented that caregivers are at a greater risk for experiencing the following problems. Are you experiencing any of them?

- Depression
- Guilt, anger, resentment & denial
- Declines in physical health
- Strained social and family relations
- Sleep disorders
- Anxiety about aging
- Neglect of self and others
- Less marital closeness
- Work absenteeism and job loss
- Loss of income
- More negative attitude toward care recipient
- Loss of free time
- Abuse of self and others
- Sleeplessness
- Exhaustion
- Loss of enjoyable activities
- Loss of friendships
- Feeling overwhelmed

Research has shown that caregivers can reduce some of these problems by using respite services along with asking for help from others. Looking out for your own well-being is similar to being a passenger on an airplane and being told that in case of an emergency you should place the oxygen mask on yourself first before caring for someone else. Why? Because you can be a more effective caregiver if you are healthy, happy and alive.

We know from research that family caregivers provide 80% of all the caregiving in the U.S. But we need to be careful in applying the label of "hero" if we end up encouraging caregivers to exceed their capabilities and sacrifice other highly valued and important parts of their lives. Depressed, divorced, and burned-out heroes will not be able to be caregivers for long. The truly heroic caregivers are those who do the very best that they can providing care, but they also recognize and value their own well-being, families and relationships and find a way to balance these competing goals.
In order to help you better understand how your caregiving is impacting some important aspects of your life, you can answer the questions on the next page (Caregiver Burden Inventory: Modified). Printed with author's permission. These 24 questions were developed by gerontological researchers Mark Novak and Carol Guest (1989).

This burden scale measures the impact of caregiving on the caregiver’s flexibility with time, physical health, social relationships, emotional well-being and life course development issues. We have used this scale in our research studies and believe that it is very helpful in identifying problem areas. We reworded two of the items (#17 & #18) to make the questions more relevant to all caregivers, which is why we call it a "modified" scale (Caserta, Lund & Wright, 1996). After each question, choose a response that describes your feelings or views. If you have a total score above 36 when you add up the points, you are very likely to be at risk for burning out as a caregiver because you indicated a response of "sometimes" on half of the items. "Burning out" means that you are feeling overwhelmed and this may lead to experiencing declines in your physical and mental health.

When scores are near or above 36 we strongly suggest that you use respite and other services. (Please remember that earlier we suggested all caregivers use respite, especially long before caregiving becomes stressful.) Before we show you another list of questions to help you evaluate your situation, we recommend that you seriously look at any item on the burden scale where your answer was scored as a 3 or 4 ("quite frequently" or "nearly always"). If you have a 3 or 4 as an answer, you need to give careful thought about why you scored so high on the question and see if you can find a way to reduce the stress you are experiencing. Some of the statements are worded to describe extreme situations and they may not match your feelings.

This scale can help you identify specific aspects of your life and situation that might need immediate attention. Also, you might consider contacting your doctor or a mental health professional to get their advice. Near the end of this brochure is a list of national and local resources that you can contact for more information and help.

We want you to take better care of yourself. Before you can do that, you need to be aware of how your life is being impacted by your caregiving.

BURDEN SCALE (Next Page)

For each item circle a number in the columns to the right that represent how often the statement describes your feelings.
<table>
<thead>
<tr>
<th>Items</th>
<th>Never</th>
<th>Rarely</th>
<th>Sometimes</th>
<th>Quite Frequently</th>
<th>Nearly Always</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Time Dependency Items</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1. He/she needs my help to perform many daily tasks</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>2. He/she is dependent on me</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>3. I have to watch him/her constantly</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>4. I have to help him/her with many basic functions</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>5. I don't have a minute's break from his/her chores</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td><strong>Development Items</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6. I feel that I am missing out on life</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>7. I wish I could escape from this situation</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>8. My social life has suffered</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>9. I feel emotionally drained due to caring for him/her</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>10. I expected that things would be different at this point in my life</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td><strong>Physical Health Items</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>11. I'm not getting enough sleep</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>12. My health has suffered</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>13. Caregiving has made me physically sick</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
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<tr>
<td>14. I'm physically tired</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
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<tr>
<td><strong>Social Relationships Items</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>15. I don't get along with other family members as well as I used to</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>16. My caregiving efforts aren't appreciated by others in my family</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>17. I've had problems with my marriage (or other significant relation)</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>18. I don't get along as well as I used to with others</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>19. I feel resentful of other relatives who could but do not help</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td><strong>Emotional Health Items</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>20. I feel embarrassed over his/her behavior</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>21. I feel ashamed of him/her</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>22. I resent him/her</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>23. I feel uncomfortable when I have friends over</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>24. I feel angry about my interactions with him/her</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
</tbody>
</table>

**Total Score = ____ (0-96)**
Caregiving presents many challenges and difficulties but being a caregiver can be rewarding as well. Other researchers have developed a set of questions that can help to identify how much satisfaction family members are experiencing by being caregivers (Tarlow et al., 2004).

Please read the following 9 questions or statements and select answers that best fit your views. Just like on the previous list of questions, you can add up all the points and learn more about your overall satisfaction from being a caregiver. We hope that you will have a score of 36 or higher, because having some positive feelings will help you be a better caregiver. Unlike the other scale, a high score on this scale indicates a positive situation. A score of 36 or higher means that you see some positive aspects related to all nine items.

Also please look at each statement carefully to learn more about where you get the greatest satisfaction or might want to get more. If you presently do not have much satisfaction, try to identify ways to enhance it. Sometimes professional help is needed through counseling or simply asking service professionals for their advice. Please do not hesitate to ask others for help.

Again, please review the list of agencies and organizations near the end of this brochure and contact them for more information and help.

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**Satisfaction Scale (Next Page)**

For each of the following questions circle a number to the right that represents how much you agree or disagree with these statements.
### Positive Aspects of Caregiving

<table>
<thead>
<tr>
<th>Providing help to this person has...</th>
<th>Disagree a Lot</th>
<th>Disagree a Little</th>
<th>Neither Agree nor Disagree</th>
<th>Agree a Little</th>
<th>Agree a Lot</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Made me feel more useful</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>5</td>
</tr>
<tr>
<td>2. Made me feel good about myself</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>5</td>
</tr>
<tr>
<td>3. Made me feel needed</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>5</td>
</tr>
<tr>
<td>4. Made me feel appreciated</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>5</td>
</tr>
<tr>
<td>5. Made me feel important</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>5</td>
</tr>
<tr>
<td>6. Made me feel strong and confident</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>5</td>
</tr>
<tr>
<td>7. Enabled me to appreciate life more</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>5</td>
</tr>
<tr>
<td>8. Enabled me to develop a more positive attitude toward life</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>5</td>
</tr>
<tr>
<td>9. Strengthened my relationships with others</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>5</td>
</tr>
</tbody>
</table>

**Total Score =** (9-45)

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Are you still doing things that you enjoy and give you pleasure?

"The primary reason for participating in an activity is to experience the pleasure of doing it."

— Jitka M. Zgola, O.T.
In our national study of nearly 900 caregivers, we found that having respite time was their most desired and needed service.

It is in everyone's best interest to encourage and help most caregivers do a bit less rather than more caregiving to restore a more appropriate balance in their lives. Sharing the daily tasks of caregiving with family members, neighbors, friends and professionals does not mean that the primary caregiver is weak, unloving, selfish, or failing to meet family obligations. Seeking and obtaining help shows maturity, commitment, concern, self-respect and an informed, positive and balanced understanding of the realities of caregiving. We hope that you are among those who are willing to ask for and seek appropriate help from others.

Caregivers often need many different kinds of help, including education, skill training, counseling, emotional, spiritual, social and financial support, legal, home maintenance and transportation services. There is a growing need for care managers, gerontological service specialists and other new types of professionals to help caregivers find their way through a complex system and arrange for an appropriate package of services to fit their specific needs, which also change over time. Many of the organizations listed near the end of this brochure can help you find the services you need.

There is one service, however, that stands out among most caregivers because it has the potential to improve or at least preserve the quality of their daily lives. In our national study of nearly 900 caregivers, we found that having respite time was their most desired and needed service (Caserta, Lund, Wright & Redburn, 1987). Another study in Michigan revealed that service professionals also identified respite as the most needed service for caregivers (Shope et al., 1993). Other studies highlighting the need for greater use of respite are increasing each year (Family Caregiver Alliance, 2006; Zarit, Stephens & Townsend, 1998).

Respite simply means having some "time away" from the responsibility of providing care. It is an opportunity for caregivers to do just a bit less, have others share the tasks and achieve some well earned balance in their lives. Respite time can be obtained at home or through a variety of adult day centers. It can range from one or two hours at a time to several days each week.

The following quotes from caregivers show some of the diverse needs that respite can help meet.

• "If only I could get a few moments of alone time, it would help me be more patient."
• "I can't take a bath, fix dinner or make a phone call without interruption. I have no privacy."
• "I need to take care of myself personally and to do things not possible when my husband is around."
• "I need time to run errands and spend more time at home alone."
• "I need to be able to work and be with other people."
• "I miss being with my friends."
• "I wish our family could take a short vacation like we did before."
• "I would like to return to work, at least part-time."
• "I wish I could go to a movie with my friends."
• "I want to play golf again or simply to take a walk."
• "I gave up most of my hobbies but would like time to do some of them again."
WHAT TYPES OF RESPITE SERVICES ARE THERE?

The United States passed the Lifespan Respite Care Act of 2006 in order to expand and improve respite care services for caregivers in every state. Respite care is seen as an important national issue.

Adult Day Services Often Provide Stimulating Activities for Clients

Although not every community has a full range of respite services available, there are several types of services. The most common form is in-home respite, usually offered through home health agencies where professionally trained persons make home visits and attend to the needs of the family member who needs care while the caregiver can leave the home.

Another common type of respite is adult day service, usually provided at a community-based site such as care centers, nursing homes, churches, assisted living facilities, senior centers and hospitals. Adult day service respite often offers the advantages of establishing regular, sufficient and relatively affordable opportunities for respite. Sometimes community or health service providers offer overnight and multiple days of respite.

This type of respite can allow families to maintain family vacations and other group activities that far too many caregivers discontinue. Remember, families need respite just as individual caregivers do, in order to preserve quality family life.

With these types of respite services becoming more available, most caregivers should have opportunities to use them and make them a regular part of a daily or weekly routine.

Another, often overlooked, form of respite is informal respite. Many friends and relatives often say, "Please let me know how I can help." Far too often, caregivers never accept these offers. Yet, having friends or relatives come to your home or take your loved one to their homes is an ideal way to share caregiving responsibilities.

You need and deserve this respite time. Rather than feeling guilty, you should allow others the opportunity to be helpful – to you and your loved one. Using respite services is one of the most effective ways to enhance the quality of life for caregivers and persons with dementia. Respite can give you a chance to help you and your loved one at the same time.

Don't delay...seek assistance today.

"Engaging in meaningful activities is important to all of us as we age; it's no different for those who are cognitively or physically impaired."

– Dale A. Lund, PhD
Getting the Most Out of Respite

Plan Ahead to Ensure That Your Respite Time Is Well Spent

Research has shown that respite needs to be used by caregivers at least two days per week, regularly, and in blocks of time to be effective (Zarit, Stephens, Townsend & Greene, 1998). However, using respite services does not automatically result in wonderful benefits to all caregivers. Early research showed mixed results about the benefits of respite help. In fact, our national research study revealed that the 36% of caregivers who were using respite services did not have lower levels of burden than non-users. More recent studies, however, are concluding that respite helps most caregivers and care-recipients but there are important suggestions that caregivers need to follow in order for respite to be MOST EFFECTIVE.

First, caregivers need to consider using respite services much earlier in their caregiving careers. Far too many caregivers try to do nearly everything on their own, without assistance, for as long as they can. Many studies have shown that respite is most helpful to caregivers when they use it before they become exhausted, isolated and overwhelmed by their responsibilities (LaSasso & Johnson, 2002; Gottlieb & Johnson, 2000). Burden levels can be high among many caregivers who use respite because they waited too long to get help. Time away from caregiving, even short periods, is needed to help maintain friendships, social activities, health and overall balance in one’s life. Once these essential features are lost, they are difficult to regain.

Second, caregivers need to have sufficient and regular amounts of respite time. Again, research has revealed that infrequent and irregular use of respite reduces its effectiveness. Caregivers need to be able to anticipate and look forward to some kind of regular or scheduled times when they know they will have at least short breaks free of the constant responsibilities of providing care. In one of our studies, 25% of the caregivers said that they did not have enough respite time. Temporary and only short-term use of respite does not provide the help that most caregivers need (Zarit, et al., 1998).

Third, caregivers need to give careful thought to how they want to spend their respite time and make the necessary arrangements in advance to ensure that they do these things while they have respite. This sounds like common sense, but it is surprising how many caregivers do not use their free time consistent with their needs and intentions. Our research team examined the ways caregivers used their respite time and was surprised to learn that 46% of them reported that they were only “somewhat” or “not very satisfied” with how they had used their respite time. We tried to identify specific activities that might lead to better results and found that the most dissatisfied caregivers spent more time than they wanted doing housework, eating, shopping, and working. Those who were most satisfied with how they had spent their respite time did things that they had wanted and planned to do (Lund, Utz, Caserta, & Wright, 2009).

Rather than suggesting one or two specific activities for respite time, it appears to be much more important that caregivers give careful thought to what they most want to do and do it when they have respite. One of the dissatisfied caregivers said, “I need to organize my time better to do hobbies.” Those who fail to plan ahead often end up “wasting time” by watching too much television or doing nothing.
If you want to go to lunch or a movie with friends it requires advance planning to make sure that it happens. When the respite time is over, it is best if you feel like the respite break was meaningful or purposeful. Some caregivers wanted to clean, work in the yard, read, be with friends or even sleep. Respite was most effective for these caregivers when they fulfilled their needs and plans.

Good use of respite time does not just happen, it requires a thoughtful review of needs, some planning and self-respect. You must plan ahead to make sure that you spend your respite time doing things that you miss the most. This is your chance to restore some balance in your life. Wouldn’t your loved one want you to do this? You may even need to be a bit selfish to make sure you spend some time enhancing your own life. One useful way to plan ahead is to take some time to set a concrete attainable goal for how you want to spend your respite time. In our research we discovered that caregivers who spent their respite time doing the things that they desired had more favorable outcomes. (Please see the goal setting box below to help you accomplish this.)

**Fourth**, respite is most effective when it is used along with other kinds of assistance. Respite is the most wanted and needed service for caregivers but it should be used along with other services or assistance and not stand alone as a single strategy (Gallagher-Thompson & Coon, 2007). Caregivers need education, emotional and social support, and a sense of belonging with others. When respite is part of a more complete package of help, caregivers are better able to meet the many needs that they have to be well, happy and enjoy daily life (ARCH National Respite Network, 2007; Feinberg, Wolkwitz & Goldstein, 2006).

Caregivers are well advised to seek help and suggestions from professionals and friends and especially from those who are very experienced in caregiving to become more aware of their own needs and what services are available. Please remember, use respite early in your caregiving career, use it regularly, engage in meaningful activities during respite time and use it along with other services.

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**USE GOAL SETTING TO MAXIMIZE YOUR LIMITED RESPITE TIME**

On a piece of paper write down 1 to 3 specific goals of what you would like to accomplish during your respite time next week. Your goals can be anything (e.g., sleeping, cleaning, going to lunch with a friend), but they must be attainable during the respite time that is available to you. We recommend that you choose activities that:

- you have **always enjoyed** doing
- you **did before** you became a caregiver but may not have been doing lately
- have **religious or spiritual** meaning to you
- may **improve satisfaction** with caregiving
- may **reduce the stress** you might feel from caregiving
- may **increase your respite time**
- encourage you to **use your respite more regularly**

At the end of the week, ask yourself whether you accomplished each of the goals and also whether you are satisfied with your effort at accomplishing each goal. Be honest with yourself, and use this information to revise your goals for the following week's respite time. Caregivers who use their respite time to do what they intended to do feel less burden and more satisfaction with their caregiving role.
### National Resources

<table>
<thead>
<tr>
<th>Organization</th>
<th>Website</th>
<th>Phone</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alzheimer’s Association</td>
<td><a href="http://www.alz.org">www.alz.org</a></td>
<td>1-800-272-3900</td>
</tr>
<tr>
<td>National Institute on Aging (NIA)</td>
<td><a href="http://www.nia.nih.gov">http://www.nia.nih.gov</a></td>
<td>1-800-222-2225</td>
</tr>
<tr>
<td>AARP</td>
<td><a href="http://www.aarp.org">www.aarp.org</a></td>
<td>1-888-OUR-AARP, 1-888-687-2277, Spanish speakers 1-877-627-3350</td>
</tr>
<tr>
<td>Alzheimer’s Disease Education and Referral Center</td>
<td><a href="http://www.nia.nih.gov/alzheimers">http://www.nia.nih.gov/alzheimers</a></td>
<td>1-800-438-4380</td>
</tr>
<tr>
<td>The Brookdale Foundation Group</td>
<td><a href="http://www.brookdalefoundation.org">http://www.brookdalefoundation.org</a></td>
<td>1-212-308-7355</td>
</tr>
<tr>
<td>Easter Seals</td>
<td><a href="http://www.easterseals.com">http://www.easterseals.com</a></td>
<td>1-800-221-6827</td>
</tr>
<tr>
<td>ElderCare Locator</td>
<td><a href="http://www.eldercare.gov">http://www.eldercare.gov</a></td>
<td>1-800-677-1116</td>
</tr>
<tr>
<td>National Alliance for Caregiving</td>
<td><a href="http://www.caregiving.org">www.caregiving.org</a></td>
<td>1-310-718-8444</td>
</tr>
<tr>
<td>Family Caregiver Alliance</td>
<td><a href="http://www.caregiver.org">http://www.caregiver.org</a></td>
<td>1-800-445-8106</td>
</tr>
<tr>
<td>National Association of Professional Geriatric Care Managers</td>
<td><a href="http://www.caremanager.org">http://www.caremanager.org</a></td>
<td>1-520-881-8008</td>
</tr>
<tr>
<td>National Family Caregivers Association</td>
<td><a href="http://www.thefamilycaregivers.org">http://www.thefamilycaregivers.org</a></td>
<td>1-800-896-3650</td>
</tr>
<tr>
<td>National Adult Day Services Association (NADSA)</td>
<td><a href="http://www.nadsa.org">http://www.nadsa.org</a></td>
<td>1-800-558-5301</td>
</tr>
<tr>
<td>Senior Corps (Senior Companions Program)</td>
<td><a href="http://www.seniorcorps.gov">http://www.seniorcorps.gov</a></td>
<td>1-202-606-5000</td>
</tr>
<tr>
<td>Rosalynn Carter Institution for Caregiving</td>
<td><a href="http://www.rosalynnncarter.org">http://www.rosalynnncarter.org</a></td>
<td>1-229-928-1234</td>
</tr>
</tbody>
</table>

### Local Resources

Nearly every town, community and city has services available to help caregivers. We suggest that you use your local telephone directory, the internet and in most areas you can call **211** for information and referral sources. You can identify and contact some of the following for help and be sure to ask about available non-profit organization:

- State unit on aging
- Area agencies on aging
- Alzheimer’s Association local chapter
- City and county social services
- Respite service provider
- Home health agencies
- Gerontology and Geriatric programs at colleges and universities

### International Resources

Alzheimer’s Disease International  
[www.alz.co.uk](http://www.alz.co.uk)  
+44 20 7620 3011

### State of California and Utah Resources

Utah Coalition for Caregiver Support  
[www.caregivers.utah.gov](http://www.caregivers.utah.gov)

California Caregiver Resource Centers  
[www.cacrc.org](http://www.cacrc.org)
The Office of Research Development (ORD)
The Office of Research Development has been established at CSUSB with funds made available by the Extramural Associates Research Development Award (EARDA) through the National Institutes of Health (NIH). The goal of ORD is to enhance faculty and student research at CSUSB, particularly in health related, biomedical and behavioral research. The ORD will facilitate faculty in their development of their research into a research program and help them develop external grant proposals. Further information about the services provided by the ORD can be found at the ORD website: http://www.ord.csusb.edu. The ORD can be reached by telephone (909) 537-3383 or via email at lgonzaga@csusb.edu or bhaddock@csusb.edu.

Research Infrastructure in Minority Institutions Grant (RIMI)
Research Infrastructure in Minority Institutions Grant was established at CSUSB with funds from the National Institutes of Health (NIH) – sponsored National Center on Minority Health and Health Disparities. The RIMI grant was designed to help minority-serving universities build innovative research programs aimed at reducing health inequalities among racial and ethnic groups and among people living below the poverty line. The RIMI grant at CSUSB seeks to: 1. Promote and accelerate faculty research in health disparities through mentorship and training from nationally recognized health disparities scientists; 2. Provide a link between health disparities-related community organizations and university members; 3. Advance the number of CSUSB students seeking graduate careers in health disparities; 4. Disseminate CSUSB faculty research and scholarship in the area of health disparities. Further information about RIMI can be found at the RIMI website: http://rimi.csusb.edu by telephone (909) 537-3635 or via email at hdinfo@csusb.edu. RIMI funds were used to support the work of CSUSB faculty to revise and update this brochure.

Community University Partnerships (CUP):
Community University Partnerships is a campus-wide initiative to build and advance partnerships in San Bernardino and Riverside Counties. We provide facilitation and support of community partnerships development, faculty fellowships to support service learning courses, and faculty mini-grants to support community-based participatory research. Contact information: Dr. Diane Podolske, Main office: (909) 537-5793 or at their website: partnerships.csusb.edu. CUP funds were used to support the printing of this brochure and also to translate the brochure into Spanish.

Your Future
We hope that this brochure will help you better understand your current caregiving situation and that you will do a better job of taking care of your own life. Many professionals, organizations and service providers are available to give you more information and offer help. Remember, it is important for you to enhance the quality of your life, and the lives of your family and loved ones.
REFERENCES


Articles on Caregiving Written by the Authors of the Brochure
(These publications are available upon request - 909-337-3748)


Dr. Michael Caserta, is a professor in the Gerontology Interdisciplinary Program and holds the Robert L. and Joyce T. Rice Presidential Endowed Chair in Healthy Aging within the University of Utah’s College of Nursing. He is a faculty associate of the University's Center on Aging and the Hartford Center of Excellence in Geriatric Nursing. He has published widely in the area of family caregiving and has been a co-investigator on research projects related to both caregiving and respite care funded by the Alzheimer’s Association, the National Center for Nursing Research, and Administration on Aging.

Dr. Chris Lindfelt came to the University from the non-profit sector. He is trained as a clinical psychologist however for the last several years he has functioned in an executive leadership role with a number of health care organizations both here and in Colorado. Most recently, Dr. Lindfelt led the Alzheimer’s Association as Regional Director for the Inland Empire. His research interests are varied. He is passionate about brain health, the aging population, caregiving, as well as current trends in long-term care and healthcare administration.

Dr. Dale Lund is Professor and Chair of Sociology at CSUSB (formerly with the University of Utah) and has conducted national and local studies on family caregiving issues since 1983. These studies identified the most common and difficult problems and satisfactions caregivers experience, their use of services and the potential of respite to be able to continue many of their lifelong interests. He served on the Executive Board of the Utah Alzheimer’s Association Chapter including two years as President and on the Strategic Planning Committee for the national Alzheimer’s Association. His caregiving research has been funded by the national Alzheimer’s Association, National Center for Nursing Research, Administration on Aging and the Bureau of Health Professions. He has published numerous articles on caregiving and made keynote presentations for the Administration on Aging, National Lifespan Respite Coalition, several universities and local Alzheimer’s Association Chapters.

Dr. Julian Montoro-Rodriguez is Professor of Medical Sociology at CSUSB since 2007. During the past few years his research was aimed at examining the interrelations between formal and informal support systems and optimal adaptation and adjustment to developmental changes for older adults. His recent work attempts to identify those conditions under which older adults and their families would increase community-based service use, in particular the implications of cultural inequalities (race, gender, socioeconomic status) as generators of distress for older adults.

Dr. Herb Shon is an Assistant Professor of Social Work at CSU, San Bernardino and a licensed clinical social worker. His main research interest is developing theory-driven, culture-specific strategies to decrease barriers to service use among under-served populations (particularly caregivers of people with Alzheimer’s disease). Dr. Shon has worked in direct services, development, and administration of Program of All-inclusive Care for the Elderly (P.A.C.E.), Adult Day Health Care, and Alzheimer’s Day Care and Resource Center programs, and is an education and training consultant to, and volunteer at, the California Southland Chapter of the Alzheimer’s Association.

Dr. Rebecca L. Utz is an assistant professor in the Department of Sociology and a faculty associate with the Center on Aging and the Institute for Public and International Affairs at the University of Utah. She teaches courses in research methods, epidemiology, and medical sociology. She has published in the areas of bereavement, aging families, and obesity.

Dr. Scott D. Wright is Director and Associate Professor of the Gerontology Interdisciplinary Program at the University of Utah. His research interests and publications are in the domains of family caregiving, environmental risk and impact in an aging society, and the role of technology in an aging society. Dr. Wright also served on the Executive Board of the Utah Chapter of the Alzheimer’s Association.
Every day is another chance to discover more about yourself—what are your strengths and limitations as a caregiver; what do you really like as a family member or friend; what are you learning about your own ability to express love, patience, and caring; and what are you discovering about your priorities in life?

— Leonard Felder, Ph.D.