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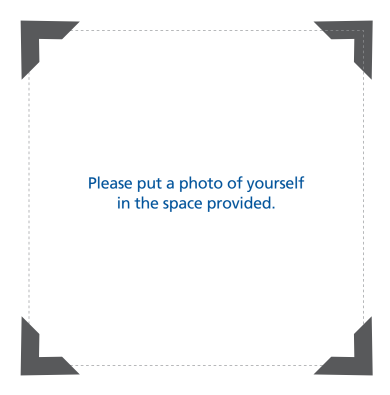
# All about me



This is a booklet about a person living with Alzheimer's disease or other dementia.

Name:

Alzheimer Society



The Alzheimer Society is Canada's leading nationwide health charity for people living with Alzheimer's disease and other dementias. Active in communities right across Canada, the Society

- Offers information, support and education programs for people with dementia, their families and caregivers
- Funds research to find a cure and improve the care of people with dementia
- Promotes public education and awareness of Alzheimer's disease and other dementias to ensure people know where to turn for help
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For more information, contact your local Alzheimer Society or visit our website at www.alzheimer.ca.

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### Introduction

This booklet is all about you, a person living with Alzheimer's disease or other dementia.

Although you have a form of dementia, you are still the same person you have always been. This booklet is designed to focus on the positive: what you are good at rather than what is no longer possible.

You and your primary caregiver\* know what makes you feel comfortable better than anyone. By answering the questions in this booklet, you will have a record of what makes you content and at ease that can be used when your primary caregiver cannot be with you and others need to provide care and support. Anyone can use this booklet to give you the best day possible now and as the disease progresses.

The first section of this booklet is designed to help someone new to supporting you get to know you better. It will also suggest conversation topics that may make you feel more at ease and contribute to more enjoyable times together.

Other sections of this booklet allow you and your caregiver to outline your usual habits: your daily routines, your likes and dislikes and what makes you enjoy each day. This information will help new caregivers maintain the routines that give you a sense of security, comfort and pleasure.

When completing this booklet, always keep in mind the main purpose: to give as clear a picture as possible of you to help others provide care when the person who usually supports you is unavailable.

To help others provide effective care, keep this book in an easy-to-find location. You and your caregivers can review it from time to time to note changes and plan for the future. There are replacement pages at the back that you can use to make any updates or changes.



<sup>\*</sup>The term "caregiver" is used throughout this booklet to mean anyone who supports you.

0	Contact information
	This booklet contains information about:
	Some of the information is provided by:
	Names, phone numbers, email addresses of significant people in my life (family, friends, neighbours):
	Name:
	Relationship:
	Phone number:
	Email address:
	Name:
0	Relationship:
<b>\_</b> /	Phone number:
	Email address:
	Name:
	Relationship:
	Phone number:
	Email address:
	Name:
	Relationship:
	Phone number:

Email address:\_\_\_\_\_

Other important numbers
Family doctor (name, phone number, address):
Ambulance:
Police:
Fire:
Poison Control:
Local Alzheimer Society:
Home-care services:
Spiritual or faith leader:
Other:
Medical information
Other than having dementia, are there other medical issues that the caregiver should know
about?
Please provide any important information on:
Allergies:
Hearing:
Vision:
Medications (attach list, if necessary) – include dosage and frequency

()

This section is like a phoof your personality.	tograph. Try to use as much detail as you can to give readers a real sense
My personal life	
How do you like to be a	ddressed? (e.g. nickname, Mr., Mrs., Miss, first name)
When were you born?	
Where?	
	ongstanding relationship(s) with
	nce(year)
Where have you lived?_	
_	
-	
	nere they are now living)
Are they involved in you	ur life now? If so, how?
Do you have any pets? I	f so, what are their names?

Date: \_\_\_

In the past What kind of jobs did you have? (e.g. homemaker, lawyer, nurse, electrician, teacher)
How do you feel about the job(s) you have done? (e.g. proud, satisfied, indifferent)
If you were asked about the major milestones in your life, what would you likely talk about? (e.g. major life events, favourite places visited)
Likes and dislikes  What makes you physically more comfortable? (e.g. always have glasses on, have a hearing aid in, daily lotion to prevent dry skin, toe spacers)
What makes you happy? (e.g. conversation topics, activities, sports, music performances, being around children/animals)
What do you dislike? (e.g. foods, activities, topics of conversation, music, smells)
What comforts you when you're upset?
What frightens you?


### A typical day

Routine is important for all of us, but can be especially helpful for a person with dementia.

Writing down your daily routine will help you see how you spend your time and help others who might be providing care.

Try to look at your care through the eyes of someone who has never met you before. Do you like to sleep in, have a bath in the evening, or go for a daily walk?

Use this section to describe regular daytime activities. Include activities you are involved in as well as your caregiver. Include anything that provides pleasure, comfort, or something you particularly don't like.

Morning (usual wake up time	). How do you start your day?
Afternoon	
Evening	
Night (usual bedtime	)

### **Enjoying each day**

A person living with dementia is just like everyone else, a whole person with likes and dislikes, opinions, values, and experience. Though some skills are lost as the disease progresses, many remain. Here are some activities that may bring you pleasure and will help you continue to live a full life while adding enjoyment to time spent with others.

What effect does it have on you	1?	
Do you play an instrument?  If yes, what kind of instrument of	☐ Yes do you play? (d	☐ No e.g. guitar, violin, clarinet)
Do you enjoy singing? What effect does it have on you	☐ Yes i?	□ No
Reading		
Do you like to read?	☐ Yes	□ No
If yes, what do you like to read?		science fiction, romance, adventure, fantasy,

•	ke watching TV? $\square$ Yes $\square$ No
If yes, w	at are your favourite shows?
Games	
-	ke to play games?
Sports	
-	nterested in sports? $\square$ Yes $\square$ No
If yes, w	at sports do you like to play or follow? (e.g. golf, hockey, tennis, skating)
Hobbies	
•	ave hobbies that you enjoy? $\square$ Yes $\square$ No
If yes, w	at kind of hobbies? (e.g. scrapbooking, crafts, photography)
Dovou	o household chores? (e.g. meal preparation, dusting, sweeping)
-	o household chores? (e.g. meal preparation, dusting, sweeping) $\square$ Yes $\square$ here any household chore you particularly enjoy?
What ot	er activities do you enjoy? (e.g. car rides, attending community programs, sitting k

Date: \_

### Help with daily living

How much help, if any, do you need with routine daily activities such as dressing, bathing or getting in and out of bed?

The chart on the next page lists typical routine daily activities. Feel free to change the chart to include activities that apply to you.

Where you are able to be completely independent, write "no help needed." When you need help, note how much help you need.

The "Useful tips" section is a good place for caregivers to note the degree of stress the activity creates and what special approaches might be helpful.

Here is a sample chart to guide you.

Activity	Useful tips	Is help needed?
Tub/shower Usual time: 8:00 a.m. Twice a week	<ul> <li>Prefer shower, don't like bath</li> <li>Enjoy music or conversation during bath time</li> <li>Give lots of time</li> <li>Respect privacy</li> </ul>	Need help in and out
Dressing	<ul> <li>Be patient</li> <li>Can button shirt, put on underwear and socks</li> <li>Need to take dirty clothes away immediately</li> </ul>	May need help from time to time
	<ul> <li>Can dress independently if clothes put on bed in right order</li> <li>Offer help tying shoe laces</li> </ul>	



Activity	Useful tips	Is help needed?
Tub/shower		
Dressing		
Dental care/dentures		
Eye care/glasses		
Lyc carc/glasses		
Hearing aid		
Hair care		
Professional style/cut		
Makeup/shave		
In/out of chair		
invoic of chair		
In/out of bed		

Activity	Useful tips	Is help needed?
On stairs		
Use of toilet		
Use of appliances		
<ul> <li>e.g. kettle, stove, electric shaver</li> </ul>		
Household tasks		
<ul> <li>e.g. sweeping, dusting, vacuuming, meal preparation, garden work</li> </ul>		
Financial		
Responsibility with money		
Walking		
Habits, usual routes, ability to be independent		
Preparing for bed		



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	nen.
וט	nner:
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_	
Sr	acks:
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Αı	ny particular likes or dislikes?
۱۸	hat assistance, if any, is required?
	utting:
_	
Us	se of cutlery:
_	
Нα	ot and cold liquids:
_	

Date: \_\_\_\_

### Regular weekly activities calendar

Use this calendar to show regular outings or appointments. You can use pencil so changes can be made every month, or a make a copy for each month.

### Month:

Sunday	Monday	Tuesday	Wednesday	Thursday	Friday	Saturday

Ö

Date: \_\_\_\_\_

0	Note: The questions in this section are designed to be answered by your primary caregiver. Your input will be valuable to give the best information possible.
	Alzheimer's disease and other dementias progress over time. As the disease progresses, your abilities will change.
	The information in this section will help anyone supporting you know what these changes are and how they affect your mood, behaviour and abilities. Your caregiver can suggest ways that help you feel content, engaged, and secure. For example – Is there a special approach that helps? Does your behaviour change only at certain times? Are there warning signs?
	Below are examples of common situations.
	The term "family member" is used to mean anyone with dementia whom you support. "He" and "she" are alternated.
	Unsafe walking (e.g. "wandering")
	Does your family member walk outside in ways that are unsafe? (e.g. will go out in winter wearing only a dressing gown)

Does your family member walk outside in ways that are unsafe? (e.g. will go out in winter wearing only a dressing gown) $\square$ Yes $\square$ No
If yes, what safety precautions do you use? (e.g. camouflaged doors, ID bracelet, regular walks with a neighbour)
Does she become upset when returned home? $\square$ Yes $\square$ No
If yes, is there a special approach to use to help her feel calm?
Is he registered with the MedicAlert $^{\otimes}$ Safely Home $^{\otimes}$ program? $\Box$ Yes $\Box$ No
If yes, what is the number of your local police station if he becomes lost?
What is his MedicAlert® Safely Home® ID bracelet #?
Night time restlessness
Does this occur? $\square$ Yes $\square$ No
If "yes," what safety precautions do you use? (e.g., nightlight, disconnecting stove or turning off water valves before retiring at night, locking closet door to prevent dressing at odd hours)

Restl	essness
Does	this occur at certain times of the day?
Wha	t helps to settle him? (e.g. a walk or a distracting activity)
Ange	er or agitation
Does	this occur at certain times? (e.g., bathing, meal time)
Wha	t usually triggers this? (e.g., rushing her, too many instructions given at once)
Whe	n anger occurs, what responses tend to be helpful?
Does	your family member suspect people of stealing from him? How do you deal with this?
Repe	tition
Whe	n he repeats himself over and over, what responses are helpful?

_	or hoarding articles
Are the	re particular places to check where your family member "stores" specific things?
Does ar	nything need to be kept out of reach? (e.g. knives, tools such as electronic drills)
Safety <sub>I</sub>	precautions
•	additional information that is important for other caregivers. (e.g. doors or cupboards ept locked, such as where toxic cleaning fluids are stored)
Are any	other safety measures being used? (e.g. alarms, GPS locating devices)
Commu	inication
-	family member has difficulty understanding and following instructions, what do you do anything that helps?
To learr the top	n more about Communication, please read the Alzheimer Society's <u>Information sheet</u> c ic.
	re any other areas of concern and/or tips for care that comfort, reassure, support the

Alzheimer's disease follows a number of stages. While these stages can be somewhat predictable, the course of the disease will vary from person to person. Changes in physical condition, such as flu, pneumonia, infection, or constipation can often result in changes in mood and behaviour. By noting changes, the person who supports you may be able to determine a pattern and prevent a situation from getting worse. It is particularly important to keep a record when medications are used. Recording these items in a journal will help caregivers when they are talking with your doctor. Caregivers should take this "All about me" booklet to your appointments.

Anyone providing care can use these pages to record all the events in a particular day.

Here is a sample journal.

Date	Comments
Mar. 30/12	To bed 8:30 p.m up again 2:00 a.m. Wandered through house, could not settle down. Did not recognize me.
Mar. 31/12	Another night with no sleep!
Apr. 1/12	Still won't sleep. Now dozing all day. Doctor's appointment April 4/12.
Apr. 5/12	On new medication for an infection. Slept till 6:00 a.m. I'm trying to keep her awake during the day. I think things are getting better.

Date: \_\_\_\_\_

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# Alzheimer Society

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