



# Your Conversation Starter Guide

How to talk about what matters to you and have a say in your health care.

Institute for  
**Healthcare**  
Improvement

the conversation project

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We can't plan for everything. But we can talk about what is most important — in our life, and in our health care — with those who matter most.

Talking with the important people in our life can bring us closer together. It also helps us create the foundation of a care plan that's right for us — a plan that will be available when the need arises.

The Conversation Project wants to help everyone talk about their wishes for care through the end of life, so those wishes can be understood and respected. We created this guide to help you start a conversation (and keep talking) so you can have a say in your health care — today and tomorrow.

It's also important to choose what's known as a health care proxy, or health care advocate — someone who would make health care decisions on your behalf if you became unable to voice those decisions yourself. Visit our [Guide to Choosing a Health Care Proxy](#) for guidance on picking a proxy.

If you are completing this document on a computer, first save it to your desktop with a name you can easily find again. Then open your saved document and type in your answers. (Otherwise, what you type will not be saved.) Completing it on your computer will create a digital document that

you can easily share with others.

~~This document does not seek to provide legal advice~~

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We'll help you take it step by step. You can take your time! There's no need to say everything that matters in one conversation — you can start talking, then keep talking. It's all about what works best for you.

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STEP 1

# Think About What Matters to You



To get ready to talk about what matters to you and your wishes for care through the end of life, it's helpful to gather your thoughts as a first step. You don't need to have the conversation just yet. Here are some helpful ways to think about what matters to you and prepare for your conversation.

**1** What does a good day look like for you?

In each statement below, mark the place on the line that is closest to how you think or believe about each statement now. There are no "right" or "wrong" answers — your answers are about what *you* think for you.

**SOME IDEAS** Is it time with family or friends? Enjoying favorite everyday activities? What do you need to enjoy a good life — through the end of life?

**2** What or who supports you during difficult times?

In the space below, write down what or who supports you during difficult times.

**SOME IDEAS** Your faith, culture, family, friends, pets

Try finishing this sentence:

What matters to me through the end of my life is...

In the space below, write down what matters to you through the end of your life.

**SOME IDEAS** Being able to recognize my children; being independent; being able to spend time with the ones I love

That's your "what matters to me" statement.

Sharing it with people you trust could be a big help if they need to communicate with your health care team one day. They may need to share what's important to you and what you need to be able to have a good day. They also may need to decide what type of treatment you'd want to receive. Completing this guide will help you refine what you want them to know about what matters to you.



STEP 2

# Plan Your Talk

Having a say in your health care is more likely if you share how you feel about certain situations that could arise now, in the future, and toward the end of life.



For each statement below, mark the place on the line that is closest to what you think or believe about each statement now. There are no "right" or "wrong" choices — your answers are about what works for you.

As a patient, I'd like to know...



Only the basics about my condition and my treatment

All the details about my condition and my treatment

When there is a medical decision to be made, I would like. ...



My health care team to do what they think is best

To have a say in every health care decision

What are your concerns about medical treatments?



I worry that I worry that I won't get I'll get too enough care much care

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If I am diagnosed with a serious illness that could shorten my life, I would prefer to...



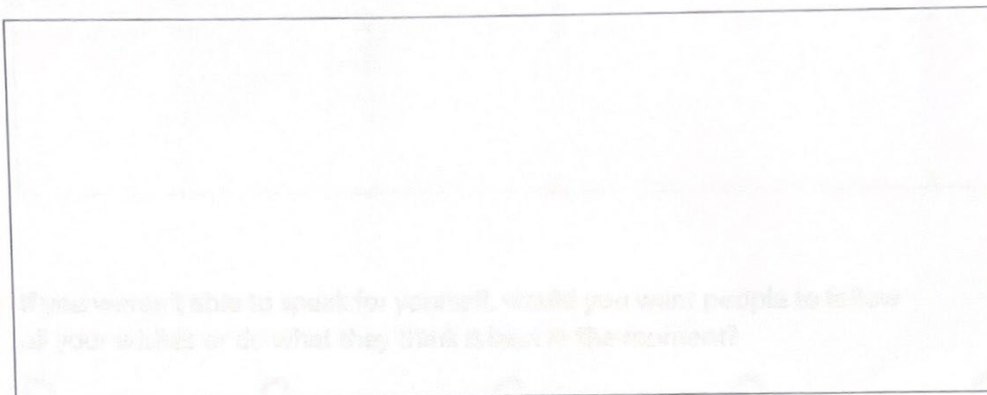
Not know how quickly

Understand how quickly

it is progressing or my doctor's best estimation for how long I have to live

it is progressing and my doctor's best estimation for how long I have to live

Any other notes you want to add?



If you were seriously ill or near the end of your life, how much medical treatment would you feel was right for you? O\_\_\_\_\_O



I would want to try every available treatment to extend my life, even if it's uncomfortable

I would not want to try treatments that impact my quality of life in order to extend my life

Where do you prefer to be toward the end of life?



I strongly prefer to spend my last days in a health care facility (hospital, assisted living, or nursing facility)

I strongly prefer to spend my last days at home

Now, look at your previous answers. What do you notice about the kind of health care you said is right for you?



If you weren't able to speak for yourself, would you want people to follow all your wishes or do what they think is best in the moment?

○ ..... ○ ..... ○ ..... ○ ..... ○

I want the people I trust to do exactly what I've said, even if it makes them uncomfortable

I want the people I trust to do what brings them peace, even if it's different from what I've said

When it comes to sharing information about my health with others...

○ ..... ○ ..... ○ ..... ○ ..... ○

I don't want those close to me to know all the details about my health

I am comfortable with those close to me knowing all the details about my health

When I die...

○ ..... ○ ..... ○ ..... ○ ..... ○

I want to be alone    I want to be with other people

What specific information would you want (or not want) shared with certain trusted people?

The screenshot shows a survey question: "What specific information would you want (or not want) shared with certain trusted people?" Below the question are two radio button options: "I don't know" and "I want to share". The "I want to share" option is selected. The text is faint and partially obscured by a watermark.

Look at your previous answers. What are the most important things for your friends, family, and health care team to understand about what matters most to you through the end of life?

The screenshot shows a survey question: "What are the most important things for your friends, family, and health care team to understand about what matters most to you through the end of life?" Below the question are several radio button options: "At the end of life", "At the time of diagnosis", "At the time of death", "At the time of recovery", "Other", and "I don't know". The "At the end of life" option is selected. The text is faint and partially obscured by a watermark.



### STEP 3

## Start Talking

How much do the people who matter to you know about what matters most to you? There may be some things they already know, and other things that you need to tell them. Sometimes we might think others know how we feel, but they don't. Conversations help make what we think and how we feel as clear as possible.

Who needs to know what matters to you in your health care?

Check all that apply:

- (O) Parent(s) Trusted friend(s)
- (C) Spouse/partner(s) Doctor(s)
- (C) Chosen family member(s) Nurse practitioner/nurse(s) o Adult child/children Social worker
- (O) Faith leader (minister, priest, C) Other: \_\_\_\_\_  
- rabbi, imam, etc.)

Where would you feel comfortable talking?

- (O) At the kitchen table
- (C) At a favorite restaurant
- (C) In the car
- On a walk
- Video chat or phone call
- At my place of worship
- C) Other: \_\_\_\_\_

The Conversation Project uses the saying, "It always seems too soon, until it's too late."

When will you start this conversation?

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You've gathered your thoughts, written down your ideas, and picked your trusted people. Now, how do you begin a conversation?

This list doesn't cover everything, but here are some things you can say to start talking.

- "I need your help with something "
- "Can you and I have a conversation about \_\_\_\_\_?"
- "I was thinking about what happened to \_\_\_\_\_ and it made me realize\_\_\_\_\_"
- "Even though I'm OK right now, I'm worried that \_\_\_\_\_and I want to be prepared. Can we talk about some things that matter to me?"
- "Will you help me think about my future?"
- "I heard about the Conversation Project and answered some of their questions about things that matter to me when it comes to my care through the end of life.  
I'd like to talk to you about it."
- "When\_\_\_\_\_died, do you think their wishes and priorities were respected toward the end of their life?"

Here is a list of some other things you may want to cover when you talk.

- Do you have any worries about your health?
- What do you need to address to feel more prepared (examples: finances, property, legal documents, relationships, health care situations)?

- Do you have any fears, concerns, or mistrust about where or how you receive health care?
  - Who do you want (or not want) to be involved in your health care?
  - When you look ahead to the future, are there important events or dates you hope you're there for?
  - Are there kinds of treatment you would want or not want (examples: resuscitation attempts, ventilation, feeding tube)?
  - If your health condition changed, when would it be OK with you to shift from trying to cure an illness to trying to enjoy the end of life as much as possible?
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## Tips for your talk

Imagine the conversation in your mind first. You can even write a letter that explains your values about the kind of care that works for you to figure out words that feel comfortable for you to use.

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- You can also consider having a practice conversation, so you feel as prepared as possible to have a "real" conversation.
- You don't have to talk about everything or talk to everyone in the first conversation. In fact, we suggest you keep talking over time!
- Be patient. Some people are nervous or may need time to get ready to talk. Every time you start a conversation, it helps you come closer to making your wishes fully known. Keep trying.
- You don't have to lead the whole conversation; it's important to also listen to what the other person says so you can build trust.
- Nothing you say is permanent. You can always change your mind as things change in the future.
- You may find out during these conversations that you and your trusted people disagree. That's OK (no judgment!). The important thing is that you're talking now and to keep talking — so you're prepared in case your health changes.
- You can share this guide, with or without your thoughts included, with your trusted people.

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## STEP 4

# Keep Talking

Now that you've started the conversation, keep going! Talk to more people who may have a say in your health care. The more you talk, the more people you are close to will know what matters to you. And that makes it more likely that you'll get the kind of health care you want — now and through the end of life. Here are some things you can think about to keep the conversation going.

- 5 When would be a good time to talk again?

**SOME IDEAS** It's a good idea to have another conversation when life changes happen, such as the birth of a baby, when family and friends are together for a holiday or visit, before a trip, or when a health issue is getting harder to manage.

What might you want to repeat or explain again, so you're sure your trusted people understand what's important to you?

- 6 Who do you want to talk to next time? Are there people (such as family members who may disagree) who should hear things from you at the same time?

- 7 What do you want to make sure to ask or talk about next time?

# What to do next

Now, it's a good idea to record your conversation with an important legal document to be sure your choices are followed. This is called an advance directive. It has two parts.

## 1. Your Health Care Proxy

This is the part of the advance directive where you name the person you have chosen to make health care decisions on your behalf, if needed, as well as an alternate if your first choice is unavailable. As explained in this guide, be sure to have a conversation — and keep talking — with these people to be sure they understand what matters to you. You can find more information and suggestions in our [Guide to Choosing a Health Care Proxy](#).

## 2. Your Living Will

This is the part of the advance directive where you describe your preferences and wishes for your health care if you cannot speak for yourself. These are many of the same things that you have thought about and discussed throughout this guide.

Every state and most countries have their own advance directive forms. In the United States, the NHPKO (National Hospice and Palliative Care Organization) can help you find the right forms in your state ([nhpco.org/advancedirective](http://nhpco.org/advancedirective)).

It's important to share your advance directive with more than your proxy alone. For example, if you pick an adult child to be your proxy and have other children, they should all be aware of what matters to you in your health care and know who you have chosen as your proxy. Talk to anyone who can help you have a say in your care through the end of life and provide copies of your advance directive to anyone who may need them. If you want tips on talking about what matters to you with your health care team, visit our [Guide for Talking with a Health Care Team](#).

### Learn more and share

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