Experiencing good NHS care: what you can expect

Information for the public
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About this information

This information explains the advice about improving the experience of care for people using adult NHS services that is set out in NICE clinical guidance 138. The information applies to people using the NHS in England and Wales.

Does this information apply to me?

Yes, if you are using adult NHS services.

No, if you are using NHS services for mental health (NICE has produced separate information for the public about this; see http://publications.nice.org.uk/IFP136).

This information does not cover carers’ experiences of NHS services.

Why NICE has written this guidance

Over the past few years, there has been an increasing focus on putting patients at the heart of the NHS, and especially on involving them fully in decisions about their care. An important development was the publication of the NHS Constitution (www.dh.gov.uk/en/DH_132961), which
sets out the rights and responsibilities of both patients and healthcare professionals. However, there is still more that can be done to make sure that all NHS patients have the best possible experience of care. This information describes what you should expect when you receive care in the NHS.

The guidance is relevant both to healthcare professionals and to other staff you may come into contact with when using NHS services. Healthcare professionals include people such as doctors, nurses, ambulance staff, dentists, dietitians, healthcare assistants, midwives, occupational therapists, pharmacists and physiotherapists. Chaplains, domestic staff, porters and receptionists are examples of other staff you may come across.

NICE has also produced a ‘quality standard’ made up of 14 statements describing high-quality care for patients in the NHS in England. The quality standard, including information summarising the statements, is available at http://guidance.nice.org.uk/QS15.

Treating you as an individual

Healthcare professionals and other people you come into contact with in the NHS should treat you with respect, kindness, dignity, compassion, understanding, courtesy and honesty. They should also respect your confidentiality, and should never talk about you in your presence without including you in the conversation.

All healthcare professionals who are directly involved in your care should introduce themselves and explain what their role is. Any students or other people present at consultations or meetings should also be introduced, and you should be able to decide whether you are happy for them to stay or would prefer them to leave.

Your condition and how it affects you

Your healthcare professional should treat you as an individual and make an effort to understand how your condition is affecting you. This involves listening to and respecting your views. They should talk with you about whether your family or work situation, or other aspects of your life, might be affecting your health and/or your ability to obtain treatment and to look after yourself. They should also ask you about your medical history and previous experiences of healthcare. They should talk with you about your mood, ask you whether you have any worries or concerns, and offer support if you need it.
Your individual circumstances

Healthcare services and care offered to you should reflect your preferences and your particular needs and circumstances, which should be assessed regularly.

Everyone should have equal access to NHS services\(^1\). If you have any physical or learning disabilities, sight or hearing problems or difficulties with reading, understanding or speaking English, you should receive appropriate support so that you can participate fully in consultations and decisions about your care. Healthcare professionals should not make judgements or assumptions about you based on your appearance or any other personal characteristics.

\(^1\)In agreement with the Equality Act 2010, which lists nine 'protected characteristics': age, disability, gender reassignment, marriage and civil partnership, pregnancy and maternity, race, religion or belief, sex and sexual orientation. For more details see www.homeoffice.gov.uk/equalities/equality-act

Help with basic needs

Making sure that a person's basic needs are met is a vital part of good-quality care. You should get the support you need to be as independent as possible, and you should get help to carry out everyday tasks if necessary, particularly if you are in hospital.

Personal care

If you need help with personal care (for example, relating to continence, personal hygiene and comfort), you should be asked regularly about what help you need. You should get help when you need it, and your privacy should be respected.

Food and drink

When you are in hospital, healthcare professionals or other staff should make sure that you have enough food and drink, which is placed so that you can reach it easily. They should encourage you to eat, and if necessary help you to do this.

Pain

If you are in pain, healthcare professionals should ask you regularly about this. They should offer
you pain relief and adjust the dose as needed.

**Medicines for long-term conditions**

If you are already taking medicines for a long-term condition when you are admitted to hospital, your doctor or nurse may discuss with you whether you want and are able to carry on being responsible for taking these medicines (rather than staff looking after them).

**Involving you in your care**

The NHS Constitution states that ‘You have the right to be involved in discussions and decisions about your healthcare, and to be given information to enable you to do this.’ The NICE guidance goes into more detail about this, as described below.

**Talking with healthcare professionals**

When you talk with a healthcare professional, they should make sure that the surroundings are suitable and that your privacy is respected, particularly if discussing sensitive, personal issues. They should ask you what you would prefer to be called, and use this name when talking to you. They should also do what they can to make you feel at ease – for example, sitting at the same level as you when talking to you. If you have had an examination and/or you are in hospital, healthcare professionals should give you enough time to get dressed or cover up before talking with you.

You should be asked whether you would like to be accompanied at consultations by a family member or friend, and whether you want to take notes and/or an audio recording to refer to afterwards. You should have access to a patient advocate (someone who helps you put your views across) if needed.

Healthcare professionals and other staff should bear in mind that accents and dialects (both yours and theirs) can affect understanding, and talk with you about this if it could be a problem. You should also feel able to raise this yourself.

If you have difficulties in speaking or understanding English, an interpreter should be provided to help you.

Your healthcare professional should talk with you about your condition and care, and about any concerns you might have. They should explain things clearly to you, including giving the meaning of any unfamiliar words. They may ask you questions to encourage discussion.
At the end of the discussion, the healthcare professional should check that you understand what has been said, and summarise the most important information.

**Information and shared decision making**

Healthcare professionals should give you all the information you need so that you can be fully involved in decisions about your care and the management of your condition.

**Involving family members and carers**

When you first visit a healthcare professional, they should check whether you want your partner, family members and/or carers to be involved in decisions about your care. If you agree, information should be shared with them. If for any reason you are unable to say whether you agree (for example, if you are unconscious), family members and carers should be kept involved and informed, but healthcare professionals should still respect your confidentiality.

**Type and style of information**

Information should be provided at the outset and throughout your care. All information you receive should take account of your preferences, and should be clear, consistent and based on the best available evidence. It should include details about your condition, treatment options, where you will be seen, who will be involved in your care, and the expected waiting times for any appointments, tests and treatments. You should be kept fully informed about the likely duration of any delays.

You should be given both oral and written information, and should have the option of getting information in different formats (such as large print, Braille, different languages, pictures and symbols), if this will help your understanding. This includes information about appointments.

You should be offered copies of letters about your care between healthcare professionals (for example, from a hospital doctor to your GP). These should be legible and, if possible, written using everyday language that you understand. Healthcare professionals should answer any questions you have about these letters.

**Making decisions about tests and treatment**

The information you are given will help you to be fully involved in and to make decisions about your treatment and care, with the support of your healthcare professionals.
When a healthcare professional offers you any test or treatment, they should explain clearly why it is being offered and the options available. You should be told about all treatment options, even if these are not available in your local area.

Your healthcare professional should ask you about your personal needs and preferences, and what you hope the test or treatment will achieve. They should provide information about the risks, benefits and consequences of the test or treatment, and make sure that you understand this information. If you want to, you should also be able to discuss the likely outcomes of any proposed treatments and your prospects for recovery.

You should have plenty of time to discuss possible tests and treatments, and be able to ask as many questions as you want. If there isn't enough time for this you should be offered another consultation.

In cases where you have a choice of treatments or tests and there is no clear 'best' treatment, your healthcare professional may offer you one or more 'tools' to help you make an informed choice. This might be a website or DVD that takes you through the various options, including information on expected benefits, risks and likely outcomes. In some cases you may be offered counselling or similar support.

Healthcare professionals should respect and support you in your choice of treatments, including if you decide not to have a test or treatment, even if your views are different from theirs. You can ask for a second opinion and should be told how to go about getting this if needed.

You should be given as much time as you need to make decisions about tests or treatments, and to talk things through with family and carers if you want to. Healthcare professionals should talk with you about what is important to you, and check that any decision you make fits with this.

Questions you might like to ask about your care

- Please tell me more about my condition and possible treatments.
- Please give me more details about possible tests.
- Why have you decided to offer me this particular treatment, and what will it involve?
- What are the benefits, risks and likely outcomes of this treatment?
Are there any other treatment options?

What will happen if I choose not to have the treatment?

Giving your consent to treatment and care

All treatment and care should be given with your informed consent. If you are not able to make decisions about your care, healthcare professionals have a duty to talk to your family or carers unless you have specifically asked them not to. Healthcare professionals in England should follow the Department of Health's advice on consent (see www.dh.gov.uk/en/DH_103643) and the code of practice for the Mental Capacity Act. Information about the Act and consent issues is available from www.nhs.uk/CarersDirect/moneyandlegal/legal. In Wales healthcare professionals should follow advice on consent from the Welsh Government (www.wales.nhs.uk/consent).

If you have made an 'advance decision' (known as a 'living will' in the past) in which you have already given instructions about any treatments that you do not wish to have, your healthcare professionals have a legal obligation to take this into account.

Information about looking after yourself

You (and your family and carers if appropriate) should be given any information you need to look after yourself. This includes information about taking medicines correctly and using any equipment you might need.

Additional information and support

Your healthcare professionals should tell you about where you can find additional reliable high-quality information and support, such as from national and local support groups, networks and information services (including websites). You should also be told about other healthcare services and social services (for example, services to help quit smoking) that are available, and encouraged and supported to use any that might be helpful for you.

Patient education programmes

If you have a long-term condition (for example, diabetes or kidney disease), you may be offered the chance to take part in an education programme, if this would be suitable for you. This usually involves going to group sessions, run by specially trained people, at which you learn more about
your condition and ways of coping with it. An education programme might also be offered as part of a rehabilitation programme (for example, cardiac rehabilitation) to help your recovery.

**Your relationships with healthcare professionals**

**Your healthcare team**

You may be able to see the same healthcare professional throughout a course of treatment, but it is more likely that you will see several members of a healthcare team. The roles and responsibilities of the various team members should be explained to you.

**Feedback and complaints**

You should be encouraged to give feedback about your care. Healthcare professionals should respond to this feedback. You should be provided with information about complaints procedures if necessary.

**Sharing of information**

Everyone involved in your care should have an understanding of your history, your condition, and any tests and treatments you have had.

Sometimes you may need to see different healthcare professionals at different places – for example, your GP and a hospital doctor, or staff in different clinics if you are being treated for more than one condition. These healthcare professionals should share relevant information with each other, and work together so that you receive the best care with the least possible delay or disruption for you. If necessary, and if you agree, your information may also be shared with social care professionals.

You (and your family members and/or carers if appropriate) should be given information about what to do and who to contact in different situations, such as 'out of hours' or in an emergency.

**More information**

The organisations below can provide more information and support for patients. NICE is not responsible for the quality or accuracy of any information or advice provided by these organisations.
Accreditation