Getting social care services when your child has additional needs

Information for parents of disabled children in England
Introduction

Local authorities have a legal responsibility to help families with disabled children. Many children have additional needs and disabilities, and some are more severely affected than others. Some disabled children and their parents will need practical support both inside the home and outside it.

This guide explains your right to get the support you need from your local authority through departments like social services, sometimes known as ‘children with disabilities teams’, ‘children’s’ or ‘social care services’.

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What is social care?

Social care is a term that generally describes all forms of personal care and other practical assistance for children, young people and adults who need extra support. This can include support provided via your local health service. Health bodies and local authorities are expected to work together to meet the health needs of disabled children. However, the focus of this guide is to provide advice about the rights of families to support from their local authority.

For more information about health services visit our website: www.cafamily.org.uk/health
Duties of social services

Social services departments have a general duty under Section 17 (10) of the Children Act 1989 to safeguard and promote the interests of ‘children in need,’ and to promote their upbringing by their families. The law recognises disabled children as being in need.

Children in need are defined as children who are under 18 years of age and:

- are ‘unlikely to achieve or maintain, or to have the opportunity of achieving or maintaining a reasonable standard of health or development without the provision of services by a local authority’, or
- whose ‘health or development is likely to be significantly impaired or further impaired without the provision of such services by a local authority’, or
- are ‘disabled’.

Section 17 (11) of the Children’s Act 1989 states that a child is disabled if they:

- are blind
- deaf, or
- non-verbal, or
- suffer from a ‘mental disorder of any kind’, or
- are ‘substantially and permanently handicapped by illness, injury or congenital deformity, or such other disability as may be prescribed’.
A child in need is entitled to an assessment from the social services department. The assessment is the start of the process to decide if services are needed. It is an opportunity for you to tell a professional about your child and family’s needs. An assessment is important because it can lead to a number of services being provided, like practical help in the home and short breaks. See page 10.

**If your child is being assessed for an Education, Health and Care plan**

Your local authority should ask you if you get any help from social services and, if necessary, make sure the social services team carry out a social care assessment, which should also consider the needs of parents and other family members. Regulations 3–10 of the Children and Families Act 2014 state that:

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**Search for the Children Act 1989 at the website below and click on section 17:**

[www.legislation.gov.uk](http://www.legislation.gov.uk)
Social care, help and support should be given to the child and family as soon as a need is identified and not wait until the completion of an EHC plan needs assessment.

For more information about getting extra help for your child at school or college, and Education, Health and Care plans, please visit our website or call our freephone helpline to speak to one of our education advisers:

☎ 0808 808 3555
🔗 www.cafamily.org.uk/education

Up to 18 years of age, your child’s needs are assessed under the Children Act 1989. The Care Act 2014 came into force on 1 April 2015 and replaced the existing system of support for adults. The changes mainly apply to disabled adults aged 18 or over and their carers. The Act place a clear duty on the local authority to ‘promote the wellbeing of disabled adults and their carers.

The Care Act also includes new rights for families with a child who may need support to prepare for the transition to adult services, and young carers. The Act makes clear that services arranged before a child’s 18th birthday to meet their assessed needs can continue after their 18th birthday until adult services are in place. See page 28.
What services can I expect for my child and family?

Services for disabled children under 18 are available under Section 2 of the Chronically Sick and Disabled Persons Act 1970, and under the Children Act 1989.

The Chronically Sick and Disabled Persons Act sets out what kind of help should be provided. Under this Act, the duty to provide services is to the individual disabled child and does not extend to other members of the family. These services are:

- practical assistance in the home, like help with the personal care of your child, for example help with getting in and out of bed
- equipment for a recreational need, like a TV, radio or computer
- leisure facilities (this could mean outings or a placement at a day centre), or
- education facilities (this could mean home-based education or funding for the personal care requirements of students so they can study)
- travel and other assistance, like travel to and from a day centre
- home adaptations and disabled facilities, for example fitting handrails or hoists
- holidays
- meals
- telephone equipment.

The Children Act 1989 sets out a range of support services, which should be available. This includes the right to permanent or temporary residential accommodation, if your child needs it. It forms the legal basis for residential short breaks. If your child needs this service, then it should be provided. If there are no suitable facilities locally, your local authority can look outside its own area.

There are other services listed in the Children Act 1989. Here are some examples:

- occupational, social, cultural or recreational activities
- home help
- assistance to enable your child and family to have a holiday
- advice, guidance or counselling
- travel assistance.

Under the Children Act 1989, local authorities have a general duty to make a range of services available to help children in the area. Importantly, the Act allows social services to provide help that will benefit other family members, like siblings and other carers.
Short breaks
The Breaks for Carers of Disabled Children Regulations 2011 requires local authorities in England to set out a range of services that will help parents of disabled children have time away from their caring responsibilities. The range of services must include:

- day time care in the homes of disabled children or elsewhere
- overnight care in the homes of disabled children or elsewhere
- educational or leisure activities for disabled children outside their homes or elsewhere
- services available to assist carers in the evenings, at weekends and during the school holidays.

Local authorities must have prepared a statement for carer, which sets out the range of services above, eligibility criteria, and how these services are designed to meet the needs of carers in their area. In addition, the statement must be published and made available on their website, reviewed, and consider the views of local people.

Some short break schemes may be described as ‘universal’, which means they are available to all children and you don’t need an assessment to access them.

ℹ️
For more information about rights to short breaks and how to challenge decisions you’re unhappy with, see our factsheet, *Short breaks – how you and your family can benefit* available free to parents who contact our freephone helpline, and free to download:

📞 0808 808 3555
✉️ helpline@cafamily.org.uk
🔗 www.cafamily.org.uk/short-breaks-factsheet
How do I access services?

An assessment is the process social services use to gather information about your child and family, so they can make a decision about what help you may need. The process for carrying out an assessment is contained within statutory guidance, *Working Together to Safeguard Children*.

This guidance tells local authorities what they must do to meet their legal responsibilities to families with disabled children. It’s up to each local authority to decide how much detail is needed in each assessment, but the maximum time frame for any assessment is 45 working days from the date they get a referral.
Urgent assessments

If you need help urgently, you can ask for services to be put in place straight away without waiting for the outcome of an assessment. The *Working Together* guidance also states that:

‘Whatever the timescale for assessment, where particular needs are identified at any stage of the assessment, social workers should not wait until the assessment reaches a conclusion before commissioning services to support the child and their family’.

The guidance also states that a child is given:

‘the support they need before a problem escalates’.

Local arrangements for assessments must be published. The social worker must make clear to children and families how the assessment will be carried out, and when they can expect a decision on the next steps. Further information may also be available on your council’s local offer website page.

A local authority social worker should make a decision about the type of response needed within one working day of a referral being received. They must also acknowledge receipt of the referral to the person asking for the assessment. Remember, if services are needed urgently they should be provided before the assessment is finished.

As mentioned earlier the maximum time frame for the assessment to finish should be no longer than 45 working days from the point of referral. If an assessment isn’t completed within 45 working days then the social worker should record the reasons for exceeding the time limit.

An outline of the process and time scales can found on pages 33 – 34 of the guidance – search for *Working together to safeguard children* at:

🔗 [www.gov.uk](http://www.gov.uk)
How do I ask for an assessment?

Many social services departments have ‘children with disabilities’ teams. You should find the number in the telephone directory under the name of your local authority (LA), social services or children's services, or call our freephone helpline for information. Your local authority’s ‘local offer’ website page should tell you about support available for disabled children and young people, and how to access it.

If you’re unclear which LA you come under type ‘Apply for a needs assessment’ into the search bar at: ✏️ www.gov.uk

Your local Family Information Service can also advise. You can find it by looking on your local authority website.

Or contact our freephone helpline for information. Our helpline team has produced a template letter you can use to ask for an assessment. It’s available on our website, or contact them:

✍️ www.cafamily.org.uk/needs-assessments
☎️ 0808 808 3555
✉️ helpline@cafamily.org.uk

Getting information about the assessment

Beforehand, you should be told how the assessment will be carried out, and be given information about what services are available – not just those which are provided by social services departments (for example, a local play scheme).

What is a ‘needs-led’ assessment?

It is very important to remember that the starting point of an assessment is your child’s needs, regardless of whether services exist to meet them. The Working Together to Safeguard Children guidance emphasises the importance of the assessment being child and family centred. The approach should also be holistic, addressing the child’s needs within their family and wider community. So, as well as your child’s disability and health needs, social services should also consider other aspects of your child’s life, for example, education and religious or cultural needs.

Preparing for an assessment

It can be useful to make a list of questions before you meet, and you are entitled to have a friend or advocate there with you. The social worker may also want to speak to your child’s health visitor, doctor or school to help them get a full picture of their needs.

What to expect at an assessment

A social worker will usually come to your home to talk to you. They should ask you for information about your child, for example, sleeping patterns, eating habits, how your child communicates, what activities they enjoy and, whether you have any other children to look after.

Don’t be worried about asking for clear information about the focus of
the assessment. It is an opportunity to have a conversation about how to meet your child’s needs. But do remember that the assessment should be based on yours and your child’s needs, and not based on services already available.

**What if an assessment has been carried out in the past?**

When services are already being provided, the assessment should be reviewed regularly. But if your circumstances have changed, you can ask for a re-assessment or review in the same way you ask for an assessment (see page 14)

**What if I’m refused an assessment?**

If your child is disabled (and in need of services) you can’t legally be refused an assessment. You don’t need to have a diagnosis for your child to get an assessment or help from social services. It can sometimes take time before a diagnosis can be made for a number of reasons, but if your child needs the type of help or support described on page 10, an assessment of their needs should still be made. Consider making a complaint if this doesn’t happen – see page 34, ‘What if you’re unhappy?’ for more information.

Advice on challenging a refusal to assess your child’s needs can be found at:

.final adopt

![Photo of a man and two children.](image-url)
Early help assessments

Often referred to as a *Common Assessment Framework* (CAF), this is another way that children and families can be assessed and receive services. But it is important to note that this process does not replace the detailed assessments under the Children Act described above. The duty to assess under the Children Act remains. However, following High Court decision *R (L and P) v Warwickshire CC*, the judge ruled that the local authority could choose to assess a child under ‘Early Help’ assessment instead of carrying out more detailed Children’s Act assessment.

The CAF can be used if you, or someone who works with your child, would like them to receive extra help and benefit from co-ordinated support from more than one agency (for example education, health and housing). It will help to identify your child’s additional needs, and other workers required to support your family.

If you and your child agree, a worker will ask you both some questions to find out what help and support you might need. This information is recorded on a simple form. Once you’re satisfied with the information on the form, you’ll be given a copy of it. Based on the information you both provide, everyone who can help your child should work together to provide the support they need.

A ‘lead professional’ will be appointed who should keep you informed, listen to your views and support you. They will also co-ordinate all the services supporting your child. You should be allowed a say in who should be the lead professional.

Further information on how the CAF operates in your area should be available from your local authority and is usually on their website.
Your needs as a carer

Any assessment of your disabled child should take into account the needs of the rest of the family members, including parents and siblings, and consider their needs as a carer and capacity to continue with caring. However, the aim of a carer’s assessment is to give you a chance to tell social services about the things that could make looking after your child easier for you. This may result in getting services or direct payments to meet your own assessed needs. Examples of services that can be provided include help with driving lessons, housework and gardening.

A carer’s assessment focuses on you as a parent and your needs. Social services should discuss issues like the help you need and whether there is anyone else who helps, or if you are your child’s only carer. The carer’s assessment should also consider your wellbeing which include the following elements:

- personal dignity and respect
- physical and mental health and emotional well-being
- protection from abuse and neglect
- participation in work, education, training or recreation
- social and economic well-being
- domestic, family, and personal relationships
- suitability of living accommodation
- the individual’s contribution to society.

All carers have a right to ask for an assessment of their needs at any time, for example if your needs have changed (you may wish to take up education, training or employment) in the same way as asking for an assessment for your child.

Parents should use the Children and Families Act 2014 to ask for an assessment. The Act also gives strengthened rights to assessments for parent carers, young people and young carers regarding work, education and leisure.

An assessment for a sibling, known as a young carer, is triggered where there is an ‘appearance of need’. That means it is not necessary for the young person to request this, so any assessment of you or your disabled child should take into account any brothers and sisters.
The Carers (Recognition and Services) Act 1995 and Carers (Equal Opportunities) Act remain in force for other family members who are providing ‘substantial and regular’ care to a disabled child. For example a grandparent, aunt or uncle who provide care, but who do not have parental responsibility for the child. Substantial and regular care is usually defined as 35 hours or more care per week.

Always put your request in writing and keep a copy of your letter or email. The request doesn’t need to be detailed but should include:

- your name and address
- details of who is in your household, including any other children
- a brief description of your child’s disability
- what kind of extra help your child needs
- whether you need help urgently.

Or you can ask your GP, health visitor, community nurse, paediatrician or voluntary organisation to contact social services to ask for an assessment on your behalf.

We have sample letters you can use to ask for a carers assessment, plus more detailed information in our factsheet, Carer’s assessments, help for parent carers available free to download or by contacting our freephone helpline

📞 0808 808 3555
✉️ helpline@cafamily.org.uk
🔗 www.cafamily.org.uk/carers-assessment-factsheet

“'The aim of a carer’s assessment is to give you a chance to tell social services about the things that could make looking after your child easier for you.’”
Tips for attending meetings with social services

➢ **Be prepared:** for example, you might want to have visited a school or a residential home if you’re planning to talk about your child going there.

➢ **Keep copies of everything:** like letters or emails you’ve written about your child to the authorities, as well as those they send you. Keep everything in a file, making sure you read everything you think might be relevant before the meeting to refresh your memory.

➢ **Make notes:** it’s easy to forget something so having a few notes handy at meetings with professionals will help to make sure you cover all the points you want to make. Listen to what the professional workers have to say too and make notes of what is said.

➢ **Take someone with you:** if you have a partner, make sure you both attend any meetings with professionals. If that’s not possible, or you are a single parent, take a friend or someone from a local support network.

➢ **Don’t be frightened:** to ask if you don’t understand what’s being said; ask questions until you do.

➢ **Keep calm:** don’t lose your temper if things are going wrong. Try to give reasoned, counter arguments.

➢ **Follow up:** compare notes and draw up a summary of the main points made at the meeting, what was agreed, and what still needs to be agreed.

“ It’s easy to forget something so having a few notes handy at meetings with professionals will help make sure you cover all the points you want to make ”
After the assessment

After social services have carried out an assessment, they need to reach a decision about whether you and your family are in need of services, and which services are needed.

They may decide there is no need for services, which could result in your case being closed with no further action taken. If you disagree with this decision, you can challenge it using the local authority’s complaints procedure. See page 23 ‘When you don’t meet the eligibility criteria’, and page 34 ‘What if you’re unhappy?’ or they may decide that there is a need for services and these should be provided. The local authority will then produce a plan of services, called a ‘care plan’.

The care plan

In many local authority (LA) areas, a panel decides the package of services that may be offered. A care plan should be agreed between social services and you and your family, to meet any identified needs. The plan should give details of:

- what services will be provided
- for how long the services are needed
- what the local authority plans to achieve by providing the services
- what each person and agency is expected to do
- the date of the next review.

Importantly, the care plan should be reviewed regularly to make sure any services remain appropriate.

It is important to seek advice if your needs change or your LA has told you a service is no longer available. If you’re unhappy with the outcome of any aspect of an assessment, see page 34 ‘What if you’re unhappy?’

When should services be provided?

Services available under both the Children’s Act and the Chronically Sick and Disabled Persons Act should be provided when there is an assessed need, and services are necessary to meet those needs. But in practice, most local authorities use eligibility criteria to help them make this decision.

Eligibility criteria

There are many disabled children in an area who need help but social services have limited financial resources. Using eligibility criteria for deciding who has a ‘need’ for services is a way they can prioritise, to make sure the people most in need get help.

The criteria are different from one local authority to another, and this means if you move to a different local authority area you may no longer qualify for the same help.
For example, one of the services listed under the Chronically Sick and Disabled Person’s Act is ‘holidays’. This doesn’t mean every disabled child must be given a holiday each time they ask. There will be local eligibility criteria. It might say for instance, that holidays will normally only be given if a child has not had a holiday for five years and there is a risk of family breakdown if it’s not given.

**When an assessment shows you meet the criteria**

Once your child’s assessed needs match the local eligibility criteria, the local authority has a duty to provide or arrange services to meet those needs.

In this example, if your child had not had a holiday for five years and you could show that your family was under such stress that family breakdown was possible, there would be an obligation to fund the holiday, regardless of the local authority’s financial difficulties.

However, the local authority can take its resources into account to decide how it will actually meet the needs. It might limit the type of holiday it provides, or it might arrange for another organisation to do so. It could even apply for a charitable grant to fund the holiday.

**When you don’t meet the eligibility criteria**

If the service is not actually assessed as a need (perhaps seen as only useful), or if it is assessed as a need but does not fit the local criteria, the local authority has no actual obligation to provide or arrange the service. They should still try their best to meet the need, for example by contacting a local charity or voluntary agency for help.

If the decision is that services aren’t needed, or that you don’t fit the eligibility criteria, you should be given clear reasons. This is to help in case you want to challenge the decision using the complaints procedures (see page 34).
Beware!
It is quite common to hear statements like, ‘Our local authority no longer provides short breaks’ or, ‘We don’t do carers assessments in this local authority.’ These statements are unlawful and you should have good grounds for a complaint (see page 34).

In fact, the local authority should not put a blanket ban on any service and should always consider the needs of the individual child and family. Other families have challenged these statements in court and the courts have decided that a local authority can’t ‘unlawfully fetter its discretion’. This means they must always be prepared to consider requests that don’t fit into their eligibility criteria.

Using the holiday example from earlier, it would be illegal for a local authority to say, ‘we never give holidays to children unless they have not had one for five years’. They can say, ‘we don’t usually give holidays’ but they must always listen to any reasons you have about why you should be treated as an exception.

Cutting back on services
When commissioning or changing services for disabled children and families, the local authority must consult with local parents via their local parent carer forum. This is a group of parents and carers of disabled children who work with local authorities, education settings, health providers and others to make sure the services they plan and deliver meet the needs of disabled children and families.

If you are told a service is being cut or changed, get in touch. You can also sign up to your local forum just to receive news for families with disabled children. To find your local parent carer forum go to the ‘Who we are’ section of the National Network of Parent Carer Forum’s website:

www.nnpcf.org.uk

Decisions on closing down certain services and facilities have also been successfully challenged in the courts because of an authority’s failure to carry out a proper consultation. Other successful challenges refer to duties under human rights and disability discrimination legislation.

For further information, see Challenging cuts to short breaks services available free from our helpline, or free to download:

0808 808 3555
helpline@cafamily.org.uk
www.cafamily.org.uk/short-break-cuts-guide
Will I have to pay for services?

Local authorities do have the power to charge for services they provide under the Children Act 1989 and the Chronically Sick and Disabled Persons Act.

Each local authority will have its own charging policy and it is usually your income and savings as a parent that are taken into account. Your child's Disability Living Allowance shouldn’t be taken into account, and you should not be asked to pay more than you can afford.

When a child reaches 16 years of age, they are assessed in their own right. This means it should be their ability to pay which is taken into account and not yours.

What if I can’t afford to pay the assessed charges?

If you feel you are being charged wrongly, or at a level that is more than you can reasonably be expected to pay given your circumstances, you can ask for the charges to be reduced or waived completely. If you’re still unhappy with the amount you’ve been asked to pay you can make a formal complaint (see page 34).

If you have any further questions about charging, contact our freephone helpline.

When should I not be charged?

If you get Working Tax Credit, or Child Tax Credit (paid at a rate above the family element) or Income Support or income-related Employment and Support Allowance, you should not be charged for Children Act services. You should also not be charged for advice, information and counselling services, regardless of your financial situation.

Do I have to pay for services provided to me as a carer?

Services for carers of disabled children are normally provided under the Children Act and the same charging rules above apply.
Personal budgets and direct payments

Local authorities might offer families the option of a personal or individual budget. This is also sometimes referred to as self-directed support.

A personal budget is an allocated amount of money that the local authority considers sufficient to meet all of your child’s eligible needs.

To decide how much will be allocated some authorities use a computerised system generally referred to as a resource allocation scheme (RAS).

Because of this method, the outcome of the RAS may lead to a shortfall in necessary services and support. Such systems have been challenged in the courts and criticised for their lack of transparency and failure to address specific individual need. If offered a personal or individual budget, you should ask the council to break down the budget to show it meets your child’s assessed needs.

Families also have the right to decline the ‘indicative amount’ and the personal budget and can insist that their care package is provided by the local authority or via direct payments. The legal duty to carry out assessments and provide services for eligible, assessed needs still remains.

Once the amount of the personal budget has been agreed, you can decide how to make best use of it. You can ask someone else (like a friend, local organisation or your local authority) to manage the budget on your child’s behalf, or you might choose to organise your child’s care yourself, taking your child’s personal budget in the form of direct payments.

Families can get a personal budget as part of an Education, Health and Care (EHC) plan, and the assessment for an EHC plan may trigger a social care assessment, as described on page 12.

For further information read our factsheet on **Personal Budgets and Direct payments** available free from our helpline, or free to download:

- **0808 808 3555**
- **helpline@cafamily.org.uk**
- **www.cafamily.org.uk/personal-budgets-factsheet**

The charity KIDS has also worked with other charities to produce guidance and resources that support families to use and manage personal budgets.

For more information visit **www.kids.org.uk/learning-and-resources.**
What are direct payments?
Local authorities can give payments, instead of providing services themselves, to allow disabled people and carers to buy in and arrange the services they’ve been assessed as needing. Any payments you receive must only be used to pay for these services.

Like personal budgets, the aim is to promote the independence of parents and disabled children who would like to manage their own social care needs. If your child is under 16 years of age, direct payments will usually be made to you as their parent. When a child turns 16, they can receive payments in their own right, to allow them to buy in the services they’ve been assessed as needing.

A request for direct payments should only be refused in very limited circumstances.

How much are direct payments?
The amount you receive should be enough to allow you to meet all the costs involved in arranging the services that social services have agreed to help with. This should include any tax and national insurance you might have to pay if you employ someone, as well as the cost of a disclosure and barring service check (previously called criminal records check).

Social services will usually deduct an amount from the payments, equivalent to what you would have been charged if they’d arranged the services. Or they may make the payments in full and ask you to reimburse them any assessed charge.

Further information about direct payments
Contact a Family has a guide Getting direct payments for your disabled child available free from our helpline, or free to download:

📞 0808 808 3555
✉️ helpline@cafamily.org.uk
🌐 www.cafamily.org.uk/direct-payments-guide
Moving into adult services

Disabled people over 18 years old have their needs met by adult care and support. This means that a disabled child receiving support from children’s services will be transitioned to adult care and support when they turn 18.

In England the rights to services and support for disabled adults and their carers are covered under the Care Act. The Act was introduced in April 2015. Provisions under the Act should ensure that there is no gap in services when a young person makes this transition. There is no duty or power for a local authority to meet the child’s needs under the Care Act before they reach the age of 18. Therefore, the general rights of parents and disabled children under 18 will remain the same. Parents still have the right to request an assessment of their child’s needs, and the local authority is still under a duty to arrange support and practical assistance in meeting those recognised needs, subject to criteria.

If a child is likely to have needs when they turn 18, the local authority must carry out a ‘child’s needs assessment’ before then, to determine what these will be. The local authority will carry out the assessment if it considers it a ‘significant benefit’ to the child to do so.

These assessments will give you an idea of the help that you and your child can expect when they move into adult care and support. After the assessment, the local authority should draw up a care and support plan; this may include a personal budget or direct payments to meet some of their needs. See page 26.
Section 59 of the Care Act also requires to local authority to provide advice and information about what can be done now in order to meet or reduce the need. They should also consider measures which can delay or prevent the development of more extensive needs.

**When can you ask for an assessment?**

There is no specific age or time before the child's 18th birthday at which the assessment must take place.

Instead, the statutory guidance suggests that these assessments take place when it is easier to understand what the needs of the child and carer will be beyond the age of 18.

For children with Education, Health and Care plans, it is likely that they will take place during the transition process, from Year 9 onwards.

The local authority must also carry out a ‘child’s carer’s assessment’ where there is ‘likely need’ for support when their child turns 18, and when it is of ‘significant benefit.’

A template letter requesting an assessment in transition is on page 30. We also have a Word version on our website you can download at

If the local authority refuses to carry out this assessment then you should ask them to provide their reasons in writing, and also ask when they think it would be appropriate to do so.

A child’s carer’s assessment will look at the carer’s ability and willingness to continue caring for their child when they turn 18, the outcomes they hope to achieve (such as paid work or study) and the support they might need to do so.

The local authority should draw up a care and support plan for the carer; this may include a personal budget or direct payments to meet some of their needs (see page 26).

This guidance provides stronger legal rights to parent carers during the transition period.

**The needs of young carers turning 18**

Young carers who provide support to their disabled sibling under the age of 18 can access support under the Children and Families Act 2014.

Young carers who are approaching the age of 18 are eligible for help in their own right under the Care Act. This is regardless of the age of their sibling.

When a young carer approaches their 18th birthday, they can ask for an assessment of their needs to find out what support can be put in place to help them achieve their aspirations, for example to go to college or work.
Dear Sir/Madam,

I am writing to request a child’s needs assessment (transition assessment) for my daughter/son under section 58 of the Care Act 2014 and chapter 16 of the Care and Support Statutory Guidance.

My daughter/son/grandchild’s is [name] [date of birth] [summary of condition]

[If appropriate] My daughter/son is currently getting support from children’s services including [short breaks/other provision].

My daughter/son has an education, health and care plan and attends [xyz school/college].

I believe my daughter/son will is likely to have needs for care and support once s/he turns 18.

It would be of significant benefit to my daughter/son to have her/his needs assessed now because [NB this is important because this assessment won’t in itself entitle your child to services before age 18. The more complex her/his needs the earlier planning needs to take place]. Give reasons why the assessment should be carried out now. For example, complex needs may require earlier planning, the time it may take to plan and put in place the adult care and support, relevant family circumstances, stage of education and upcoming exams, plans to enter into work or training, plans to move out of parental home, planned medical treatment.

Please contact me at the above address or telephone number to let me know when you will be able to carry out an assessment. I look forward to hearing from you in a timely fashion as required by Care and Support Statutory Guidance. If you are unable to carry out this assessment then please provide your reasons in writing and also let me know when you think it would be appropriate to do so.

Yours faithfully,

[your name]
When a child turns 18
The Care Act is intended to prevent a gap in services when a child turns 18.
For this reason, the Act makes clear that any children’s services a child is receiving before their 18th birthday will continue after their 18th birthday, until adult care and support takes over.
The same is true of a carer receiving support from children’s services when their child is under 18. The local authority must not allow a gap in care and support when young people and carers move from children’s to adult services.
If a carer is not currently receiving services when they receive a child’s carer’s assessment, the local authority can choose – but is not legally required – to meet a carer’s needs before the child reaches the age of 18.

The rights of disabled adults and their carers
If a disabled adult has been assessed as having needs for care and support from the local authority, they will receive a care and support plan, and advice about decisions on how to meet their needs. They might have some of their needs met via direct payments (see page 26).
Under the Care Act, charging reforms were scheduled for introduction in April 2016. These have now been postponed to April 2020. In the meantime, the local authority may charge for services depending on the disabled adult’s finances. People will only be asked to pay what they can afford, and this will be decided by a financial assessment carried out by the local authority.
The Care Act provides much greater rights for carers of adults aged 18 and over. For the first time, the needs of a disabled adult’s carer will be treated in the same way as the needs of the disabled adult themselves. Carers can ask for an assessment if they ‘appear’ to have needs for support. This is a low threshold and will entitle most carers to an assessment.
The assessment will take into account the carer’s wellbeing, any outcomes they’d like to achieve, whether they are willing and able to care for the disabled adult, and whether they would like to access work, education or training.

For information visit:

对我来说，这个文档的内容主要是关于儿童和残疾人的护理服务。该法案旨在防止儿童在18岁时服务中断。法案明确指出，任何在18岁之前接受的儿童服务将在18岁后继续，直到成人护理和支援接管。同样的情况也适用于在18岁以下的孩子接受儿童服务时的看护者。地方当局不得允许在年轻人士和看护者从儿童服务到成人服务时出现照顾和支持的中断。

如果看护者在收到儿童看护者的评估后未收到服务，地方当局可以选择——但不是法律要求——在儿童达到18岁时满足看护者的需求。

《残疾人及看护者权利》
如果一个残疾成年已被评估为有护理和支援的需求，他们将收到护理和支援计划，以及关于如何满足其需求的建议。他们可能有一些需求可以通过直接支付来满足（见第26页）。

根据《护理法》，收费改革原定于2016年4月实施。这些收费改革已被推迟到2020年4月。在此期间，地方当局可以就服务的收费决定是否符合残疾成人的财务状况。人们只会被要求支付他们能负担得起的费用，这将由地方当局进行财务评估决定。

《护理法》提供了更大的权利，为18岁及以上的成人看护者。首次，一个残疾成人的看护者的需求将得到与残疾成人自己相同的方式对待。看护者可以要求评估，如果他们‘看起来’有需要支持。这是一个低门槛，将使大多数看护者能够获得评估。

评估将考虑看护者的福祉，他们想要实现的任何结果，他们是否愿意并能够照顾残疾成人，以及他们是否想要获得工作、教育或培训。

有关更多信息，请访问:

- [www.cafamily.org.uk/moving-into-adult-services](http://www.cafamily.org.uk/moving-into-adult-services)

见我们的事实文件《准备成年》免费从我们的热线获取，或免费下载:

- [0808 808 3555](tel:08088083555)
- [helpline@cafamily.org.uk](mailto:helpline@cafamily.org.uk)
- [www.cafamily.org.uk/adulthood-factsheet](http://www.cafamily.org.uk/adulthood-factsheet)
If your child can’t manage their affairs

Some people have a disability or type of illness that affects their ability to understand and make decisions for themselves. They may need help with managing their affairs and making personal welfare decisions. This may affect decisions that need to be made about their education, health, social care, money or where they live.

Most day-to-day decisions about daily living won’t be controversial, as parents and carers are expected to act in their child’s best interest. But some professionals may disagree and question the actions of the parent or carer. If you are concerned about this, seek further advice about your options in ensuring your views are being listened to.

In difficult cases, under the Mental Capacity Act, steps can be taken to ensure that a parent or carer has the authority to make key welfare decisions on behalf of the disabled person age 16 or over.

This involves applying to the Court of Protection to become a deputy, which incurs costs and legal fees. However, the Mental Capacity Act 2005 Code of Practice states that ‘deputies’ for personal welfare decisions will only be required in the most difficult cases where:

- important and necessary actions cannot be carried out without the court’s authority, or
- there is no other way of settling the matter in the best interests of the person who lacks capacity to make particular welfare decisions.

If a personal welfare decision is not controversial and no-one opposes it, it may be possible to make the decision on behalf of the person who lacks the mental capacity to make it her/himself without referring to the Court of Protection.

For further advice you can contact the Office of the Public Guardian on

The Office supports and promotes decision making for those who lack capacity:

📞 0300 456 0300

See also Special Educational needs and disability code of practice – Participating in decision making, page 20, from paragraph 1.3. And Annexe 1 of the code of practice. Search for the title at: 🌐 www.gov.uk
With regard to benefits, young people are expected to manage their entitlements from the age of 16. A few months before a child’s 16th birthday the parent will be contacted by the Department of Work and Pensions (DWP). They will arrange an appointment to establish whether the parent or carer should continue to be their child’s appointee and deal with their benefits. If they feel a young person is not capable of handling their affairs and managing benefits, then the parent will be asked to carry on doing so as their appointee.
What if you’re unhappy?

Under The Children Act 1989 ‘Representations Procedure (England) Regulations 2006’ parents and carers can complain about:

- service quality or appropriateness
- delays in decisions being made or services being put in place
- how services are delivered (or not delivered) including the way complaints are dealt with
- the amount of help given, how frequently a service is provided, any changes made to services or how much you are asked to pay
- the attitude or behaviour of staff
- how eligibility and assessment criteria are applied
- a local authority policy which impacts on you or your child
- any aspect of the assessment, reviews or care management.

Similar rights exist in respect of complaints about adult social care services.

Who deals with a complaint?

Each local authority has a ‘designated officer’ who receives all complaints, called the complaints manager. They don’t have to handle all stages of the complaint but are responsible for administering the scheme to make sure complaints are dealt with swiftly and effectively.

How quickly can I expect a complaint to be dealt with?

Time scales are summarised below but you can ask the authority to respond sooner.
**Stage 1 – local resolution**
You should bring your concerns to the attention of the person providing the services locally. The local authority should consider mediation at this stage, and all other stages.

They should make a first attempt to resolve matters within 10 working days. This can be extended by another 10 days, for example if an advocate needs to be appointed.

If the matter isn’t resolved, or if there is agreement for an investigation to take place, then the complaint should go to Stage 2.

If you wish the complaint to go to this stage you can request this orally or in writing.

**Stage 2 – Investigation**
The local authority should arrange an investigation that produces a report and a decision within 25 working days (or sometimes, in extreme circumstances, this can be extended to 65 working days). If the matter is still not resolved then you can ask for a panel to consider your complaint. The investigation will be undertaken by a nominated complaints officer.

**Stage 3 – Review Panel**
The person making the complaint can ask for the matter to go to a Review Panel within 20 working days of receiving a Stage 2 decision. This is a meeting of three independent people who will consider the complaint and make recommendations. The process of holding a Review Panel must follow certain time limits. See box on page 36.

**Unhappy with the outcome of a Review Panel?**
If, after a Review Panel has considered your complaint, the matter is still not resolved then you can take your complaint to the Local Government Ombudsman (LGO). You may be able to approach the LGO earlier – for more information see the next page.

You can find further information about complaints in government guidance *Getting the best from complaints – social care complaints and representations for children, young people and others*, by searching for the title at [www.gov.uk](http://www.gov.uk).

**Freephone helpline 0808 808 3555**
<table>
<thead>
<tr>
<th>Action</th>
<th>TIME FRAME</th>
</tr>
</thead>
<tbody>
<tr>
<td>You request a Review Panel</td>
<td>Up to 20 working days after receipt of the Stage 2 decision</td>
</tr>
<tr>
<td>Complaints manager acknowledges request</td>
<td>Within two working days</td>
</tr>
<tr>
<td>Complaints manager appoints a Chair and confirms attendees and content of the panel papers</td>
<td>Within 10 working days of your request for Review Panel</td>
</tr>
<tr>
<td>Local authority agrees the other panellists and date for Review Panel</td>
<td>Within 30 working days of your request for Review Panel</td>
</tr>
<tr>
<td>Local authority circulates panel papers</td>
<td>Within 10 working days of the date for the Review Panel</td>
</tr>
<tr>
<td>Review Panel produces its written report (including any recommendations)</td>
<td>Within five working days of the Review Panel meeting</td>
</tr>
<tr>
<td>Relevant director issues their response</td>
<td>Within 15 working days of receiving the Review Panel’s report</td>
</tr>
</tbody>
</table>
Local Government Ombudsman

If you’re not happy with the panel’s decision then there are various other options, like taking the issue up with your local authority (LA), or MP, or complaining to the respective Local Government Ombudsman (LGO).

The Ombudsman can investigate complaints against principal councils (not town, parish or community councils) and certain other bodies. By law, they must look for ‘maladministration’ by a local authority. Maladministration is a term that describes the actions of a government body that can be seen as causing an injustice. Examples include:

- delays
- incorrect action or failure to take any action
- failure to follow procedures or the law
- failure to provide information
- inadequate record-keeping
- failure to investigate
- failure to reply
- misleading or inaccurate statements
- inadequate liaison
- inadequate consultation, and
- broken promises.

The Ombudsman will decide whether the LA has done something wrong which has directly affected you and caused you an injustice. They won’t investigate a complaint against the local authority simply because you disagree with it. There are two LGOs in England that deal with complaints from different parts of the country. Details are at:

📞 0300 061 0614
➡️ www.lgo.org.uk/contactus

Complaints against your local authority

Each year the Local Government Ombudsman sends an annual review to all local authorities (LAs) about complaints made about them to the ombudsman. This is a useful way of assessing the LA’s performance in dealing with complaints. The review includes statistics but also lists a few cases that have been investigated and reported. To see your LA’s review letter visit:

➡️ www.lgo.org.uk/information-centre/councils-performance
Taking legal action

Judicial Review
If your complaint is very urgent and you can’t wait for the complaints procedure to resolve the matter, you can apply to the courts for a Judicial Review. Judicial Review is a procedure where the High Court looks at the way a decision was reached to see if it was legally correct.

You can also apply for Judicial Review if you have exhausted the complaints procedure and are still unhappy with the outcome. To do this you will need legal assistance.

If you have a low income, you may qualify for legal aid. In addition, some solicitors offer a free first interview.

Further information about help with legal costs can be found at:

[*www.gov.uk/legal-aid*]

For advice about seeking legal help visit:

[*www.cafamily.org.uk/legal-advice*]

Or contact our freephone helpline:

[*0808 808 3555*

[*helpline@cafamily.org.uk*]
Other ways to complain

Ofsted is the Office for Standards in Education, Children’s Services and Skills. They inspect and regulate services that care for children and young people, including children’s homes. They also inspect services that provide education and skills for learners of all ages. Ofsted have some powers to investigate concerns about services they inspect or regulate. For more information visit:  

🔗 www.ofsted.gov.uk/contact-us/how-complain
📞 0300 123 4666

Healthwatch England is the consumer body for users of health and care services. They have the power to raise concerns and influence the policy of health and care providers. You can contact your local Healthwatch, which may be able to offer advice and support with your complaint. For more information visit:  

🔗 www.healthwatch.co.uk/complaint

The Care Quality Commission regulates and inspects children’s homes that provide healthcare performed by a qualified healthcare professional. It also regulates and inspects home care agencies that provide services for children. For more information visit:  

🔗 www.cqc.org.uk/content/share-your-experience

Both Ofsted and the Care Quality Commission value hearing about any concerns about the care provided by services they inspect. If you're concerned about the care your child is getting, do share it with them, as other parents may have reported concerns too. It helps them build up a picture of what is going on in a local area and may trigger an investigation if parents are reporting bad practice.
Frequently Asked Questions

My social worker said I should put my disabled child on a register of children with disabilities. What does this mean?

Social services have an obligation to keep a register of children with disabilities. This isn’t the same as the child protection register and doesn’t suggest in any way that your child is at risk. You don’t have to agree to your child’s name being added to the register and it doesn’t affect your entitlement to services.

A register allows social services departments to try to plan services for disabled children more effectively in their area. It is sometimes used as a way of getting relevant information to families so you might get a newsletter as a result.

Can I use direct payments to employ my sister-in-law who lives with me to look after my son?

The 2003 Direct Payments Regulations make it that clear that a direct payment should not be used to employ a spouse, partner or close relative living in the same household as the disabled person.

However, employing a close relative may be possible if this is the only effective way to meet a child or young person’s needs. For example, if a child has complex communication needs or specific cultural needs then it may be acceptable to employ a family member who could meet those needs.

My daughter is disabled and I’m wondering if I can get a Blue Badge?

The Blue Badge Scheme is a UK arrangement of on-street parking concessions for disabled people. The badge can also be used in many European countries.

If your child has severe walking difficulties, is registered blind, has severe upper limb disabilities or receives the higher rate mobility component of Disability Living Allowance, you may be eligible. If you are the parent of a child who is less than three years old, you may apply for a Blue Badge for your child if they have a specific medical condition which means that they either:

- must always be accompanied by bulky medical equipment which cannot be carried around without great difficulty, or
- need to be kept near a vehicle at all times, so that they can, if necessary, be treated in the vehicle, or quickly driven to a place where they can be treated, such as a hospital.

For more information contact your local authority or visit: www.gov.uk/apply-blue-badge.
My son has ADHD and is also autistic. Social services won’t assess my child, saying they can only help children with physical disabilities and they don’t have any suitable services

Under the Children Act 1989, a local authority has a duty to assess any child in need. See the definition of ‘children in need’ on page 4 of this guide. The definition includes a reference to someone who ‘suffers from mental disorder of any kind’. Whilst this phrase is outdated it can include a wide range of conditions, including Asperger syndrome/high-functioning autism, attention deficit hyperactivity disorder (ADHD) and attention deficit disorder (ADD) as well as impairments such as learning disability, mental illness and personality disorder. In light of this definition, it would appear that social services have acted incorrectly by refusing your request for an assessment and failing in their general duty to provide a range of services for all disabled children. But you should note that they’re also allowed to set eligibility criteria for specific services, which could mean that your child won’t qualify for a service because they don’t meet the criteria. If this is the case and you’re unhappy about their decision, you can make a complaint using the local authority complaints procedure. See page 34.

More advice written by community care lawyers on challenging a ‘refusal to assess’ decision can be found at: www.cafamily.org.uk/refusal-letter

Can my social worker advise me on which benefits my family are entitled to?

Social workers do have a duty to provide advice and guidance. They should explain to you what benefits are available and make sure you’re getting the benefits you’re entitled to.

However, most social workers are not benefits experts and they may rely on the local authority’s welfare rights service or similar advice agency to help you. They may also direct you to a voluntary organisation like the Family Fund for further help. Advice about your entitlements can be found in a range of guides from Contact a Family, available from our freephone helpline. Helpline advisers can answer benefit enquiries and refer you to our welfare rights team for specialist advice if you need it. More information, including advice about changes to benefits can be found on our website at: www.cafamily.org.uk/benefits
Due to the cuts, I’m worried that I will lose help the local authority provide with getting my daughter ready for school in the mornings. I’m disabled too and couldn’t manage without this.

Your local authority must continue to provide the service as long as you continue to meet the criteria you were assessed against. In addition, the authority can’t take its resources into account if you would be left at severe physical risk if the help wasn’t provided.

However, the local authority can take its resources into account when setting criteria for services. This means they may choose to introduce new stricter criteria, which could make it harder for you to access the same level of help. Even if they introduce new criteria they can only reduce or withdraw a service after carrying out a re-assessment of your needs, and you don’t meet the new criteria. You may still be able to challenge a decision to withdraw or reduce a service in this way by using the complaints procedure or taking legal action. Call our freephone helpline for further advice. Or you visit our website to see how you can make a complaint: [www.cafamily.org.uk/services-complaints](http://www.cafamily.org.uk/services-complaints).

Do I have a right to see my child’s records?

Under the Data Protection Act 1998 professionals and agencies have a duty not to disclose information about disabled children and their families without the consent of the ‘subject’. This applies to children as well as adults provided that, if they are under 16 years of age, ‘they have the ability to understand the choices and their consequences’.

Even though there’s a duty not to disclose without the child or young person’s consent, the public body concerned (for example, the health service or local authority) still have the discretion to allow access to the information. In most cases, parents should have no difficulty seeing their child’s records.

Guidance also says that advocates should be given access to relevant information about the person they’re representing. Our freephone helpline can give you more advice about access to personal records.

I contacted social services for help with getting a short break and accessing local activities and play schemes but I was told that this is what my child’s Disability Living Allowance should be used for. Is this correct?

Disability Living Allowance is not intended to replace funding from
the local authority where there is an assessed eligible need. You should still ask for an assessment. An assessment of your child’s needs will hopefully identify the need for short breaks and lead to services being provided. This may include the offer of direct payments, which can be used to employ someone to enable your son to attend an activity that he enjoys. If you’re told otherwise, ask for this in writing and then challenge their decision. See page 34 for advice about challenging decisions. Our guide *Holidays, Play and Leisure* has more information about your child’s rights to access play and leisure opportunities. Call the helpline for a copy or visit our website where you can download it at: www.cafamily.org.uk

I’m finding it very difficult to cope and need help at home, but I’m scared that if I contact social services they’ll take my children away.

Unfortunately, many parents feel that needing help will lead to social services taking their child away. Some even feel that a request for help is a sign of weakness or bad parenting. For this reason, we understand that some families will not seek help.

It’s important to understand that while social services have specific duties around child protection issues and protecting vulnerable children from harm, they also have duties and responsibilities to support families.

This means where possible keeping the family together.

Social services should respond to your situation by carrying out an assessment to identify your needs. If you like, you can also ask for a separate assessment as a carer. Either way, this should lead to a recognition of your needs and the practical support you’re looking for to help ease the situation at home.

Some parents get regular breaks, which allow them to spend time, either with their other children or alone, so they can recharge their batteries. Spending time away from your disabled child may also help foster a sense of independence. This may be particularly helpful with transition to adulthood, as they grow up.

If you’re still concerned about approaching social services, you may find it helpful to contact a voluntary organisation for support and advice. They may be able to support you in your request for help or tell you about other services, which aren’t linked to social services.

Contact our freephone helpline for further information:

📞 0808 808 3555
✉️ helpline@cafamily.org.uk
My husband and I are asylum seekers and care for our severely disabled daughter. We can’t claim any benefits and we’re not allowed to work. Instead, we receive asylum support from the UK Border Agency (UKBA). We’re finding it very difficult to lift our daughter in and out of the bath and onto the toilet. She also needs a wheelchair - without one we can’t take her anywhere. Are we entitled to any extra help?

The UKBA are responsible for maintaining you, your husband and your child. They are responsible for providing for a child’s ‘essential living needs’. This means the needs of ‘an ordinary child or adult with no special peculiarities or disabilities’.

Although section 122 (5) Immigration and Asylum Act 1999 states that local authorities cannot provide ‘assistance’ under section 17 of the Children Act 1989 to a child whose parents are receiving UKBA asylum support, this doesn’t prevent them from providing support to a child who is disabled. Any needs that go beyond ‘essential living needs’, such as the needs arising from a child’s disability, should be provided for by the local authority.

Under the Children Act 1989, the local authority can provide further support and services for your daughter’s needs as a disabled child. If UKBA asylum support is inadequate, you can approach your social services department for help.

It is important to seek further clarification about your rights to make sure you don’t jeopardise your status and current support. You may find, for example, that getting practical help from another source to meet another need could affect the asylum support you get from UKBA. Call our freephone helpline for further information on specialist sources of advice.

I am a student from overseas, living with my wife and disabled child. Because of our immigration status, we’ve been told we have ‘no recourse to public funds’. What does this mean?

Most people coming to stay in the UK under the immigration rules are expected to be able to house and support themselves financially, without having to rely on public funds or state support. This is what is meant by ‘no recourse to public funds’.

In reality, this means you’re not usually allowed to claim benefits like Income Support, Housing and Council Tax benefits, Disability Living Allowance and tax credits. In addition, you’re not usually allowed to apply for local authority housing.

For the purposes of immigration rules ‘public funds’ does not include NHS treatment, state education or help from social services, for example under the
Children Act 1989. If you’re not clear about your rights, it’s very important to seek specialist advice. If you don’t, it could affect your status and right to remain in the UK.

The Contact a Family freephone helpline can provide limited advice and assistance on immigration matters, including advice on rights to entitlements. But we can’t help with more complex issues like challenging a failed application for asylum or advising on deportation. If we can’t give the help you’re looking for, we should be able to suggest where to get help in your area.

Contact our freephone helpline for information about specialist immigration advice services:

📞 0808 808 3555
✉️ helpline@cafamily.org.uk

Parent Advisers can arrange for an interpreter if English is your second language.
How Contact a Family can help

Contact a Family is a UK charity that provides support and information to families with disabled children, whatever the condition or disability.

📞 Our helpline

Our freephone helpline can give advice about any aspect of raising a disabled child, including help with finances, education, emotional and practical support.

0808 808 3555   helpline@cafamily.org.uk

📚 Guides for parents

We have a range of free guides for parents, including:

- Helping your child’s sleep
- Siblings
- Fathers
- Developmental delay (explanation and developmental milestones)
- Claiming Disability Living Allowance (the main benefit for disabled children)
- Personal Independence Payment and other benefits at 16

A full list of our guides is at the link below. All our guides are free to parents who call our helpline, and are free to download.

www.cafamily.org.uk/publicationslist

0808 808 3555

This guide was written by Jesslyn Parkes
Get in contact with us
209–211 City Road, London EC1V 1JN
📞 020 7608 8700
✉️ info@cafamily.org.uk
🎥 www.cafamily.org.uk
🌐 www.facebook.com/contactafamily
🐦 www.twitter.com/contactafamily
📺 www.youtube.com/cafamily

Free helpline for parents and families:
📞 0808 808 3555 (Mon–Fri, 9.30am–5pm)
✉️ helpline@cafamily.org.uk (Access to over 200 languages)