Working together to support Young Carers and their families
Foreword

We are proud of the children and young people who are carers in Cumbria and know how much they value their caring roles.

The 2011 National Census indicated that there are 166,363 young carers, 20% up from 2001 (and widely believed to be an under-representation of the true figure; in 2010, a BBC and university of Nottingham survey suggested there could be around 700,000). We should also take into account that these latest figures may not recognise young people who care for an adult with mental health issues or substance misuse difficulties.

In Cumbria it is estimated that there are approximately 3,378 Young Carers in the County. 850 of these children are known to Young Carer Organisations who are commissioned on behalf of Cumbria County Council to provide services to young carers. We want to ensure that young carers are identified and supported and that they are not taking on inappropriate levels of care whether the care needs arise from ill health, disability, mental health needs or substance misuse. We know that if care-giving becomes excessive there can be risks to the emotional or physical well being or educational achievement and life chances of the young carer.

This Cumbrian joint Memorandum of Understanding between Cumbria County Council’s Health and Care and Children’s Services sets out the new responsibilities created by the Children and Families Act and Care Act and are placed on a local authority as a whole (not just Adult or Children’s Services) together these pieces of legislation are intended to provide a legal framework that will support local authorities to consider the needs of the whole family, deliver coordinated packages of support and protect children and young people from excessive or inappropriate caring roles.

The memorandum is a commitment that our services will work together to ensure that:

- we develop better ways of identifying where caring by children becomes inappropriate, or excessive, and that:
- we find ways to put in place the support that prevents this happening.

It provides a useful framework for staff who engage with young carers, adults, or siblings who use care service and places an emphasis on taking a ‘whole family’ approach as outlined in the Children and Families Care Act. It outlines our responsibilities and reminds us about the questions we should ask in our work with children, young people and their families.

This memorandum is aimed at all practitioners in Cumbria whether caring for adults or children, from schools, mental health services and substance misuse services, Children’s Services, Health and Care Services, voluntary sector services and other agencies providing social or educational support.

It has been adopted by Cumbria County Council Children’s Service and Health and Care Services and is endorsed by Cumbria’s Local Safeguarding Children Board.

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Working together to support Young Carers and their families

What we are aiming to do locally – a joint statement of intent

Young carers tell us that they value their caring roles and are often proud of the contribution they are able to make in their families. All too often, however, children and young people become carers because someone in their family has significant unmet care needs arising from ill health, disability, mental health needs or substance misuse. In some cases young carers have stepped into the breach, sometimes assuming a level of responsibility that no child should be expected to take on. This can have consequent knock-on effects on schooling and other key areas of their lives.

Putting People First\(^1\) emphasised that care services should be delivered in ways which sustain families, avoid the need to take on inappropriate caring roles and prevent further inappropriate caring. This policy aim, which is also reflected within the current national strategy for carers, is interdependent with the principle of integrated working.

Making it Real [2011]\(^2\) was prepared by the Think Local Act Personal Partnership [TLAP] and sets out a framework for taking forward personalised, community based support.

Positive for Youth, 2012,\(^3\) the cross-Government policy for young people aged 13-19 offers us real insights and encouragement on how we can work together in partnership to support families and improve outcomes for young people; especially, those who are vulnerable.

We have committed to working together locally. We will do this across systems, in partnership with health and local carers’ organisations and within the resources available. We will work in partnership with parents and young carers to ensure:

- Children have a sense of belonging within supportive relationships where parents feel supported in their parenting role.
- Risks to independence, safety and welfare are recognised and responded to. We ensure safety of those who are vulnerable and at risk of significant harm and do so in ways that are personalised, proportionate and risk based.
- Integrated, earlier and more effective responses to young carers and their families are adopted using approaches such as the “whole family pathway”.
- There are no “wrong doors”. Young carers are identified, assessed and their families are supported in ways that prevent excessive or inappropriate caring and support parenting roles regardless of which service is contacted first.
- No care or support package for a parent or sibling relies on excessive or inappropriate caring by a young carer to make it sustainable.
- Young carers are encouraged to have strong ambitions and good opportunities to realise their potential and to have the same access to education, career choices and broader opportunities as their peers.
- Transition to adulthood is supported. The challenges faced by young adult carers [18-24] around education, training, employment and independence are responded to.
- All young carers and their families feel empowered. Increasingly they see themselves and are seen as partners in shaping what we do.
- We learn from and build on their experience and outcomes.
Empowerment

1 Young Carers: a shared understanding
We are agreed that the term “young carer” should be taken to include children and young people under 18 who provide regular and ongoing care and emotional support to a family member who is physically or mentally ill, disabled or misuses substances.

The term does not apply to the everyday and occasional help around the home that may often be expected of or given by children in families and is part of community and family cohesion. The key features for us are that:

“caring responsibilities are important and relied upon within the family in maintaining the health, safety or day to day well-being of the person receiving support or care and/or the wider family.”

We will continue to work together to develop a shared and more detailed understanding of the different types and levels of caring in our area. Our main focus, however, will be to ensure we develop better ways of identifying where caring by children risks becoming excessive and/or inappropriate and putting in place the support that prevents this happening.

The central issues for us are recognition, adverse impact, empowerment and support, including emotional support and accountability. Timely assessment and early intervention can prevent a child undertaking inappropriate levels of care. We start from the belief that:

“a young carer becomes vulnerable when the level of care-giving and responsibility to the person in need of care becomes excessive or inappropriate for that child, risking impacting on his or her emotional or physical well being or educational achievement and life chances”

The young carers involved in the ‘Whole Family Regional Conferences’ facilitated by The Children’s Society provided powerful testimony about joint working and support services for young carers. They want to be seen as just like other children and young people. At the same time they are very clear that timely and effective support for young carers and their families can make a real difference to the impacts they experience by:

- reducing marginalisation, isolation and anxiety
- managing feelings of stigma or shame
- meeting additional needs
- keeping together as a family – being a family
- enabling them to keep up with school work
- improving school attendance and achievement
- enabling young carers to continue in education post 16 and gain employment
- recognising what it means to be a “young carer”
- responding to concerns around self identification and what happens next

2 Promoting awareness and recognition
Councils must ensure that young carers are identified and that consideration is also given as to whether they are a child in need (Section 17 Children Act 1989). Councils are required to be proactive about identifying young carers in their area and acting to help reduce their need for support through the provision of information and preventative measures (for example in schools).

We have heard key messages that:

- Without early identification young people’s disclosure tends to happen at crisis points.
- Young people appreciate professionals who give them space to build trust as well as the choice to talk, what to tell and at what pace.
- Young people’s repeated experiences of disrupted relationships with professionals may result in resignation and lack of engagement.
One in 12 young carers are caring for more than 15 hours a week, and around one in 20 misses some school because of their caring responsibilities.

There is a fairly even split between the proportion of young carer aged 10-14 (41%) and 15-17 (46%); one in eight are under the age of ten.

Young carers are 1.5 times more likely than their peers to be from Black, Asian or minority ethnic communities, and twice as likely not to speak English as their first language.

The average income for families with a young carer is £5,000 less than for families who do not have a young carer.

There is no strong evidence that young carers are more likely than their peers to come into contact with support agencies.

Young carers have lower educational achievement at GCSE (equivalent to nine grades lower overall) and are more likely than the national average to be not in education, employment or training (NEET) between the ages of 16 and 19.

We will keep local practice under review and where appropriate refine it to ensure that it:

- promotes positive images of adults with long term conditions/disabilities that encourage families to seek information, assistance and support children with caring responsibilities;
- provides appropriate and accessible information for families about services that support parenting capacity, independence and well being;
- enables access to self directed support; including direct payments to meet the needs of parents where appropriate;
- reaches out to families to offer support that avoids inappropriate forms of caring developing or continuing;
- reflects principles of partnership working with communities, in particular, the need for sensitivity to cultural needs;
- supports schools in their key role of identifying children with additional support needs and early intervention and support of young carers;
- identifies “hard to reach” groups/families and creates opportunities to meet their needs;
- recognises that care needs can vary significantly and ensures local processes offer emergency advice and support where usual care arrangements risk breaking down; and,
- engages with local young carers’ projects for early support and whole family working.

Awareness is the key to recognition. Indicators of the impact of caring on children can include:

- Problems at school, not completing homework, absenteeism, lateness and inability to take part in after school activities.
- Social isolation from other children their age, feeling that no one else can understand his or her experience.
- Lack of free time for play, sports and leisure activities.
- Emerging behavioural problems, in some cases including youth offending activity.
- Emotional impacts, such as worry, depression, self-harm.
- Physical impacts, such as tiredness, fatigue, back injury.
- Lack of aspirations and career opportunities.
- Increased independence and maturity for their age.
- Advanced life skills such as a caring attitude or being a good listener.
- Increased knowledge of disability and illness.

Not all children who have ill or disabled parents or siblings take on caring roles or do so in ways that cause difficulties. Circumstances will vary. What is important is that we work closely with the family and the young person so that reasonable steps can be taken to pre-empt likely problems and any emerging difficulties affecting well-being can be identified at an early stage.

Health and Care Services, in addition to assessing parental social care needs, have a key role in identifying young carers, as they will often be the first point of contact. At the point of assessing the cared-for person, Health and Care Services will ask whether the person they are assessing has children and, if they do, what impact they feel their situation has on them.
SIGNPOSTS [ADCS/ADASS 2011] contains valuable evidence to inform practice on working together to improve outcomes for young carers in families affected by enduring parental mental illness or substance misuse. It is a useful resource for local professionals in identifying and supporting young carers. It offers points for discussion that we can use to support progress.

3 Schools, academies and colleges

Schools, Academies and Colleges will be encouraged to identify young carers at an early stage; promote and co-ordinate their support of young carers; and, liaise with other agencies as appropriate with the outcomes we are seeking. School nurses also have a role to play here. We will encourage schools and academies to:

- have a named staff member with lead responsibility for young carers and to recognise this role within continuing professional development.
- have in place a policy to encourage practice that identifies and supports young carers such as adapting school arrangements if needed, provision for personal tutors and private discussions and access to local young carers’ projects.
- promote open communication with families that supports parenting capacity and encourages the sharing of information.
- ensure school policies such as those for enrolment, attendance, bullying, behaviour and keeping safe afford recognition to young carers.
- incorporate into individual pupil plans recognition and support for the positive aspects of the young carer’s role, as well as providing the personalised support necessary to enable young carers to attend and enjoy school.
- consider scope for school staff to adopt lead professional roles within locally agreed assessment arrangements or CAF.
- consider the role of school nurses in supporting improved health outcomes and reduce inequalities of family/child experience.

4 Promoting health and wellbeing

Health professionals are also likely to be among the people that a family turns to for help with an illness or disability. Whether they work in a hospital or community, with adults or children, they may be the only person who is able to ask the right questions to find out that a child is taking on caring responsibilities. Additionally, we will encourage GP surgeries to have registers identifying carers and young carers and consider use of e-learning resources.

Child and adolescent mental health services should be used as appropriate to support the emotional well being of young carers who are seriously troubled by their caring role. Integrated working across health, Health and Care Services, Children’s Services and third sector partners and through local partnership arrangements and the local Health and Wellbeing Board will be used to develop a strategic and operational framework that identifies young carers and their needs. This would be done with a view to:

- Promoting and sustaining healthy lifestyles and diets
- Encouraging regular exercise
- Ensuring good oral health
- Raising awareness and reducing risks of substance misuse
- Offering smoking cessation support to young carers interested in giving up.
- Raising awareness of maintaining emotional well being and reducing personal stress
- Enabling young people to assess risks about lifting and handling and provide information, advice and support to remove or reduce risk of injury as necessary
- Promoting safe procedures for control of medication that do not involve young carers.

5 Equality & diversity

As with abuse or neglect, inappropriate caring responsibilities adversely impacting on wellbeing, cannot be condoned on gender, religious or cultural grounds. We will ensure that appropriate people are readily available to provide advice on such matters. We will tackle barriers to effective communication and take up of support.
When considering translation services we will consult with families as to who could fill this role appropriately. Where appropriate and possible, bi-lingual advocates will be used and account taken of any relevant factors around faith, gender or locality. We are agreed it is not good practice to expect young carers to interpret for their families, particularly when it involves someone with an illness. We will discourage this. We expect interpreters to be used and will reinforce this in staff guidance as appropriate. There may be occasions, however, where a family express a strong preference for an adult family member to be the interpreter. Where all are in agreement and the requirements and responsibilities of the role are understood this can be considered.

We will keep under review and encourage staff awareness around gender issues and assumptions that can impact upon both male and female young carers.

6 Information for empowerment
Together with our partners, we will work towards a position where, if not already in place, information and advocacy services are available to all young carers and their families offering information, advice, advocacy, representation and support. This includes, where appropriate, peer support through local young carers’ projects or parenting groups. We will encourage local use of the following general principles when people act as advocates for young carers and/or their families:

- Advocates should be the individuals’ person of choice and can be informal as well as professional advocates. Peer advocacy may be appropriate in some situations.
- Advocates should work for the best interests of the individual and their family.
- Advocates should be alert to the potential for conflicts of interests within families and potential needs for separate advocates in some situations.
- Advocates should value and respect young carers and their families as individuals and challenge all types of unlawful discrimination.
- Advocates should work to make sure that everyone understands what is happening to them, can make their views known and exercise, where possible, appropriate choices when decisions are being made.
- Advocates should help young carers and their families to raise issues and concerns about things with which they are unhappy. This includes complaints.
- Advocates must understand requirements regarding safeguarding and know what to do if they become aware of abuse or neglect or risk of it occurring.

7 Information sharing
Effective and timely information sharing between our agencies and with our partners is critical to empowerment, the provision of early intervention and preventative work, supporting transitions and, for safeguarding and promoting the welfare of young carers. Within the framework of existing local information sharing protocols our aim is to ensure specific recognition of the position of young carers. This will cover their identification and support. Local arrangements for information sharing under this protocol will be consistent with national guidance. All practitioners should follow the seven “golden rules” that are in place:

- Remember that Data Protection legislation is not a barrier to sharing information
- Be open and honest about why, what, how and with whom information could be shared,
- Seek advice if in any doubt
- Share information with consent where appropriate
- Consider the child’s safety and welfare
- Gather and keep secure information that is necessary, proportionate, relevant, accurate, and timely
- Keep a record of decisions and what, if any, information has been shared and with whom.

8 Young Carers in transition and Young Adult Carers
The Care Act introduces new obligations to young carers “in transition” to adulthood. The Act requires councils to undertake an assessment for a young carer if it considers that she / he is likely to have needs for support after becoming 18 and that the assessment would be of
significant benefit to him / her. If a local authority decides not to undertake such an assessment it must give reasons for its refusal. This change will also provide new opportunities to improve the identification of young adult carers (aged 18 to 24).

We will build on local experience and make use of the findings of Young Carers Pathfinders and other research to deliver our commitment on transition to adulthood and for support of young adult carers. We will:

- Raise professional awareness of the risks and challenges faced by young carers around low aspirations, negative experiences of learning and support and the impacts of disadvantage and consequences of caring responsibilities on take up of education, training and employment.
- Aim to have one organisation/named professional who takes responsibility for the holistic needs of a young adult carer’s; support on transition issues, moving from dependence to independence; improving resilience and opportunities to take up education, training and employment whilst recognising needs around continuing to care.

**Assessment**

9 **Introduction**

Cumbria County Council has a statutory duty to provide an Assessment of needs for support and is extended to all young carers under the age of 18 regardless of who they care for. LA should make links between children's and adults legislation to enable LA to align the assessment of a YC with an assessment of an adult they care for (preventing inappropriate care).

The Care Act requires councils to undertake an assessment for a young carer if it considers that he / she is likely to have needs for support after becoming 18, this should provide new opportunities to improve the identification of young adult carers (18 to 24).

We are agreed that the key to ensuring better support and outcomes for young carers is effective assessment. If a referral is made about a parent with a disability, dependency or illness, agencies should always consider whether there is a child in the family who is providing personal care or practical support. In doing so, practitioners will be expected to consider, preferably within a whole family approach, the impact of the disability or illness on each child within the family; including whether any of them are or could be providing care or support that is relied upon, is impacting on wellbeing and where a review of adult care needs is indicated.

Concerns may arise in many different contexts and their nature will vary. Our local approach will make appropriate use of partnership and integrated working. For young carers and their families this includes:

- **Assessment** – ensuring all assessments are timely, transparent and proportionate within the locally agreed Assessment Framework or Early Help Assessment which is consistently understood and applied. [See Below].
- **Early intervention** – early or identification of situations before they become critical
- **Reviewing or referring for review** Health and Care Services will renew the needs of a parent where children with caring responsibilities that are relied upon within the family are identified.
- **Personalising Support** – using the potential of personalised care and self-directed support planning to meet care and support needs.
- **Recording** – making sure information is in one place with the consent of the child or parent consistent with established principles for obtaining informed consent.
- **Sharing information** – so that all agencies involved know what the issues are, what is intended and so that young carers and families do not have to repeat things to us. [See above]
- **Joint Decisions**, using, as appropriate, Team around the Child and Team Around the Family for assessments and evidence based decisions for support
- **Lead Professionals** – acting as the point of contact for young carers and their families to make connections, build trust, bring things together and help them stay that way.
- **Ensuring child safety** [See: p 15]
- **Effective professional supervision and regular reviews** – seeing assessment as a continuing process to ensure a clear direction of travel and inform future plans.
10 Empowering and proportionate

The local approach to working with families will be empowering, holistic, inclusive, proportionate, integrated, child centred, rooted in child development, focus on strengths as well as difficulties and have a clear focus outcomes. We will:

- Consider the family as a whole, acknowledge parents’ strengths, promote resilience and beware of undermining parenting capacity.
- Work with colleagues from all sectors including with the voluntary sector where appropriate.
- Ensure that the assessment process is appropriate to age and understanding and specific to their needs as a young carer.
- Recognise that families may be fearful of acknowledging children’s caring roles.
- Ascertain if the illness/disability is stable, changing or episodic.
- Maintain a focus on positive outcomes for the young person and their family when working with other departments/agencies.
- Respond to young carers’ needs for emotional support and counselling.
- Consider the family’s housing needs and access to benefits.
- Be sensitive to cultural perceptions and needs around disability, illness and caring consistent with a child’s fundamental right to a safe and secure childhood.
- Recognise there may be differences of view between children and parents about appropriate levels of care and that such differences may not be acknowledged.
- Take account of the young carers wishes regarding education, employment and recreational activities.

The resolution of any tensions requires good quality joint work between Children’s Services and Health and Care Services as well as co-operation from schools and health care workers. This work should include direct work with the young carer to understand his/her views. The young person who is a primary carer of his or her parent or sibling may have a good understanding of the family’s functioning and needs. These should be incorporated into any assessment.

This memorandum also provides a framework to ensure that any lead professional, Health and Care Services or Children’s Services, should have access to and hold multi-agency information and assess the whole family regularly. Consideration will be given to who is deemed to be an appropriate lead professional having regard to all the circumstances of the assessment.

We will encourage professionals to ask certain questions either as part of their assessment, or during professional supervision, or at review to inform judgements about what is in the “best interests” of the young carer and their family. These questions might well include:

- Is a child undertaking (or at risk of undertaking) caring tasks likely to impact on them?
- Why is a child undertaking care and support tasks that are relied upon?
- What is the impact of caring on the child’s development, health and well-being?
- What additional personalised services or support may be needed to ensure the parental care needs are met or to sustain a family unit and to prevent a child taking on or continuing to hold inappropriate caring responsibilities?
- What is the parental capacity to respond to needs? Do they need support in their parenting role or in developing their parenting capacity?
- What can be done to help the whole family or to maximise the broader support which others in the family are able to provide and to promote resilience?
- How might we build resilience and family strengths and manage risks along the way?
- Do the impacts on the child indicate that it would be appropriate to engage the locally agreed framework for assessment of Children in Need and their Families or under the Carers and Disabled Children Act 2000, Children and Families Act or Care Act?
- Are there any additional needs falling within the locally agreed Assessment Framework for Children?

Keeping the Family in Mind offers some timely reminders from children and young people for professionals coming into contact with parents with enduring mental health needs. We will encourage professional awareness of these, as appropriate, along with the principles of successful front line family services.
11 Whole family working
Care Act (Statutory Guidance 6.43 to 6.50)
A whole family approach will be embedded into local assessments. We will ensure that:

- The primary responsibility for responding to the needs of young carers derives from the person in need of care and support. This means that whichever service identifies there is a young carer in the family, whether it is children’s or adults’ social care services or health, it is responsible for referring or assessing the needs of that young carer within that family context.
- Practitioners seek advice and support where necessary from colleagues, whether it is children’s or adults’ social services or a partner agency, to support discharge of our joint and separate responsibilities towards young carers and their families.
- Practitioners are aware of the prejudices and stereotypes that may exist around cultures, and disability, or about adults who misuse drugs/alcohol or have mental health needs in terms of their parenting capacity and competence.
- Practitioners reach their conclusions on the basis of the evidence of their observation of both parents and children; including any young carers.

12 Focused on Change and Outcomes
Providing an assessment only for the child will not necessarily resolve the situation that has caused their referral. All Health and Care Services and children’s assessments should ascertain why the child is caring, the extent of the reliance and caring responsibility and what needs to change. This is essential to prevent children from undertaking inappropriate levels of care and being relied on to assume levels of responsibility which impact adversely on their own well-being.

Timely assessments of both the person who needs care and the whole family could prevent a child undertaking inappropriate levels of care in the first place. When a referral is made about a parent with a disability, substance dependency or illness, we have committed to finding out whether there is a child in the family who is providing personal care or practical help. In doing so, professionals will also be expected to consider, within a whole family approach, the impact of the disability or illness on any child within the family; including, whether any of them are or could be providing some form of care or not. Similar considerations apply if there is a child with a disability within a family.

Such assessments should not only identify regular individual personal care needs (including safeguarding), but should also consider the range of parenting, caring and family tasks that are needed when care workers are not present and mean a child is relied upon to carry them out.

13 Joint assessment
Joint assessment by Health and Care Services, child and family and health staff will be expected where this is appropriate. Access to specialist advice and support should be available as needed. Finally, we should never ignore any aspect of a situation that indicates there are concerns about children’s and/or vulnerable adults’ safety and they require protection from harm.

Safeguarding

14 Children at risk of harm
Safeguarding is part of a continuum where prevention and early intervention can help young carers and their families work through the challenges they face. Safeguarding is about keeping children safe from harm and abuse and is an important part of integrated working.

By working together in an integrated way professionals place the child at the centre of all activities and are better able to identify holistic needs earlier and improve outcomes. We accept a joint responsibility to work in partnership with others to identify and respond to any young carers who are suffering, or likely to suffer, significant harm and to protect them from this harm. We will do this in ways that keep children safe and:
focus on working together, early intervention and prevention;
reflect practice guidance;
do not stigmatise families or risk increasing the number of hidden young carers; and,
do not discourage young carers and their families from seeking information and advice, or an assessment and provision of services.

Local single and multi-agency policies and procedures set out clearly the local arrangements for safeguarding children at risk of significant harm and/or promoting their welfare. We will:

- State clearly the responsibilities of staff under local safeguarding children procedures to make referrals where children are considered to be suffering or likely to suffer significant harm and emphasise the principle that safeguarding is everyone's business.
- Ensure all staff and volunteers across all sectors have undertaken appropriate training in recognising harm, reporting concerns about a child's welfare and safety and confirming referrals they have made to children's social care within 48 hours.
- Ensure all staff and volunteers across all sectors have undertaken appropriate training in relation to mental health and substance misuse issues.
- Make sure our arrangements for young carers and their families reflect any requirements of local multi-agency and single agency policies for safeguarding children and seek inclusion as necessary.

15 Adults at risk of harm

The Vision for Health and Care Services identifies seven key principles for building up a modern system of social care. They are: prevention, personalisation, partnership, plurality, protection, productivity and people. Protection is defined as ensuring that:

‘there are sensible safeguards against risk of abuse or neglect. Risk is no longer an excuse to limit people's freedom’.

We are agreed that we have a joint leadership responsibility to:

- Ensure awareness of safeguarding adults’ policy and practice; the ability to recognise and respond to safeguarding adults’ concerns; and to promote confidence and consistency in using local multi-agency procedures by staff in across all agencies.
- Apply the agreed principles of adult safeguarding and secure consistency with local multi-agency policies and procedures in respect of adults who are vulnerable and more at risk of harm in line with the following:
  - Empowerment: presumption of person led decisions and informed consent.
  - Protection: support and representation for those in greatest need.
  - Prevention: it is better to take action before harm occurs.
  - Proportionality: proportional and least intrusive response appropriate to the risk presented.
  - Partnership: local solutions through services working with their communities.
  - Communities have a part to play in preventing, detecting and reporting abuse and neglect.
  - Accountability: accountability and transparency in delivering safeguarding; including learning from experience and outcomes

16 Local safeguarding boards

Local Safeguarding Children and Adults Boards have been made aware of the general issues surrounding young carers and the intention to adopt this Memorandum of Understanding. This has been done to ensure consistency with local multi-agency policies and procedures.

It is also intended to raise awareness of the way in which safeguarding work forms part of a continuum of locally agreed person-centred and proportionate risk-based responses. We can all use these to ensure that those adults and children at risk of harm are kept safe and their welfare is promoted.
Accountability

17 Funding responsibilities

The internal allocations of funding by the Council should not become a barrier to timely and appropriate support for young carers and their families. We recognise that disputes about where funding responsibility lies can be deeply damaging to families. They were one of the concerns voiced by families and young carers in national consultations on the National Carers Strategy. We will act to ensure that staff have a clear understanding of joint and separate responsibilities to support parenting roles, respond to needs and reduce the need for inappropriate caring by young carers. The following general principles apply to the expected whole family and joint approach to meeting needs and arranging support:

- Health and Care Services is responsible for commissioning care and support services for adults to reduce or prevent inappropriate caring responsibilities by young carers.
- Children’s Services is responsible for commissioning services to respond to specific needs of the child or young person; including, those relating to the impact of their caring role on them.
- Shared responsibility exists between us for commissioning services that would support or sustain adults in their parenting role having regard to the individual circumstances.

18 Preventing disagreements

We believe that the inclusive, whole family approach to which we are committed should mean significant disagreements between local Children’s Services and Health and Care Services will be rare. Two potential areas suggest themselves and are:

- disagreements about whether the need relates to the young carer or the adult or sibling who is supported by him or her; and/or,
- disagreements about respective responsibilities or thresholds for adults or children.

We intend to reduce the risk of disagreements by:

- ensuring that staff are appropriately trained and supported in understanding and in the exercising of joint and separate responsibilities towards young carers and those they support;
- being as clear as we can about our joint and separate responsibilities;
- ensuring young carers and parents have access to information and advocacy services to support them in the exercise of their rights; and,
- ensuring that effective arrangements for consultation, communication and feedback to young carers and those they support are available and acted upon.

How such issues are resolved is a matter for us as the Statutory Directors to determine within the context of our corporate responsibilities within the Council. The following general principles will be used to inform action and decision-making:

- Disagreements about funding responsibilities must not get in the way of responding in a timely manner to situations where it is evident that inappropriate caring responsibilities are being undertaken and relied upon.
- Disagreements about funding must not be allowed to become a problem for the young carer or the person supported and must not be argued about in front of them.
- Disagreements about responsibilities must not leave the needs of family members unmet because they seem to fall between internal administrative boundaries.
- Dispute resolution procedures relating to the joint and separate responsibilities of Statutory Directors for young carers and the people they support will be put in place.
- Both Statutory Directors have final operational responsibility for ensuring that any disagreements about funding are resolved in a reasoned, timely and appropriate manner with better outcomes for young carers being a primary consideration.
19 Learning and development
We will ensure that our programmes for learning and development reflect the need for joint and separate training to underpin the organisational, policy and practice principles adopted for working with young carers and their families.

Feedback from young carers and their families will be used to inform our programmes.

20 Local partnerships
We are agreed that successful local partnerships depend on the building of constructive relationships and a shared vision around what we are trying to do. We will use the opportunities for working together to identify key priorities for commissioning and the best use of available resources designed to secure the outcomes for well-being we have identified and agreed.

Appendix A

Working together to support young carers and their families within a whole family approach

Flow chart

For all statutory, non-statutory and independent sector services working with, or likely to come into contact with families and young carers.
Flow chart for professional when a referral is made for an adult with a disability or illness

1. **Does the adult have children?**
   - Make a note of this, including any ages.

2. **Is the adult ‘vulnerable’ and at risk of harm?**
   - Contact Safeguarding Adults officer - consider safety.
   - www.cumbria.gov.uk/adultsocialcare/stayingsafe

3. **Is the child at risk of significant harm?**
   - Contact Safeguarding Hub.
   - www.cumbrialscb.com

4. **Is there a child that may be providing care and support?**
   - As why and what needs to change to reduce or prevent inappropriate caring?

5. **Are there any other agencies and departments in contact with the family?**
   - Contact them, share information and work together.

6. **Have adult care needs been assessed?**
   - Check if an Initial Needs Assessment has been done.
   - Join up with the lead professional to ensure it takes into account parenting needs.

7. **Is caring role or responsibilities impacting or likely to impact adversely on the child’s wellbeing?**
   - Join up with Children’s Services - consider requesting an assessment (Carers Assessment or locally agreed assessment / Early Help Assessment) Refer to Young Carers service.
   - Re-assess the parents’ needs, taking into account their parenting needs.

8. **What is needed to support the whole family in ways that protective were needed, promote resilience or support parenting responsibilities?**
   - Ask the client, record it and join up with relevant agencies to provide whole family support.
   - Consider a multi-agency meeting to discuss the whole family’s needs. Agree who is providing what and regularly review the needs of the whole family.

9. **Is the condition and general situation of the adult stable or is it changing or episodic?**
   - Assessments and care packages consider how support will:
     - Meet changing or episodic conditions.
     - Include plans for crisis provision to prevent a child from taking on caring role in the future.
Flow chart for professionals when a referral is made for a child who is a young carer

1. **Is the child at risk of significant harm?**
   - Contact Safeguarding Hub.
   - [www.cumbrialscb.com](http://www.cumbrialscb.com)

2. **Is the adult ‘vulnerable’ and at risk of harm?**
   - Contact Safeguarding Adults officer - consider safety.
   - [www.cumbria.gov.uk/adultsocialcare/stayingsafe](http://www.cumbria.gov.uk/adultsocialcare/stayingsafe)

3. **Ask why the child is caring and what needs to change.**
   - A referral of a Young Carer to Children’s Services should automatically trigger a review of the needs of the person who requires care (parent/sibling/grandparent)
   - More information: [cumbria.gov.uk/healthandsocialcare/adultsocialcare](http://cumbria.gov.uk/healthandsocialcare/adultsocialcare)
   - [sicarers.org.uk](http://sicarers.org.uk)
   - [westcumbriacarers.co.uk](http://westcumbriacarers.co.uk)
   - [furnesscarers.co.uk](http://furnesscarers.co.uk)
   - [carlislecarers.com](http://carlislecarers.com)
   - [edencarers.co.uk](http://edencarers.co.uk)

4. **Is the caring role significant enough to impact on the child’s life?**
   - Refer the young person to an appropriate support service.

5. **Has a locally agreed assessment or Early Help Assessment been done.**
   - Join up with the lead professional to ensure it takes into account parenting needs.
   - Consider whether a local assessment / Early Help Assessment is appropriate.

6. **Is the child’s school aware of family situation?**
   - If appropriate (ask) contact them, signpost to education support.

7. **What can be offered to support the whole family to promote resilience and support parenting responsibilities?**
   - Join up with relevant agencies to provide an emergency plan to prevent caring in the future.
   - Consider a multi-agency meeting to discuss the whole family’s needs. (Team around the Family - TAF). Agree who is providing what and regularly review the needs of the whole family.
Appendix B

Recognised valued and supported - the current national policy context for carers

_Recognised, Valued and Supported [2010]_ set out the Coalition Government’s broad approach and priorities in England with a view to securing the best possible outcomes for carers and those they support.

The five key outcomes within the 2008 strategy continue to inform the overall framework:

- Carers will be respected as expert care partners and will have access to the integrated and personalised services they need to support them in their caring role.
- Carers will be able to have a life of their own alongside their caring role.
- Carers will be supported so that they are not forced into financial hardship by their caring role.
- Carers will be supported to stay mentally and physically well and treated with dignity.
- Children and young people will be protected from inappropriate caring and have the support they need to learn, develop and thrive and to enjoy positive childhoods.

The Coalition Government identified four key priority areas flowing from consultation responses and discussions with the Standing Commission on Carers. They are:

- Supporting those with caring responsibilities to identify themselves as carers at an early stage, recognising the value of their contribution and involving them from the outset both in designing local care provision and in planning individual care packages.
- Enabling those with caring responsibilities to fulfil their educational and employment potential.
- Personalised support both for carers and those they support, enabling them to have a family and community life.
- Supporting carers to remain mentally and physically well.

The priority areas were recognised to be overlapping and that “… addressing any one of them adequately will require attention to all of them.”

_Source: Recognised, Valued and supported: next steps for the carers strategy [2010]_
Appendix C

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