Severe mental illness and substance misuse (dual diagnosis) – community health and social care services

NICE guideline
Draft for consultation, May 2016

This guideline covers how to improve services for people aged 14 and above who have severe mental illness and misuse substances. The aim is to provide a range of coordinated services that address their wider health and social care needs as well as other issues such as employment and housing.

Who is it for?

- Health, social care, community and voluntary sector organisations offering services for people with a severe mental illness who misuse substances.

It is also relevant to:

- People aged 14 and above with a severe mental illness who misuse substances who live in the community, their families and carers and the public.

This guideline contains the draft recommendations, information about implementing the guideline, context, the guideline committee’s discussions and recommendations for research. Information about how the guideline was developed is on the guideline’s page on the NICE website. This includes the evidence reviews, the scope, and details of the committee and any declarations of interest.
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**Recommendations**

People have the right to be involved in discussions and make informed decisions about their care, as described in *your care*.

*Making decisions using NICE guidelines* explains how we use words to show the strength (or certainty) of our recommendations, and has information about prescribing medicines (including off-label use), professional guidelines, standards and laws (including on consent and mental capacity), and safeguarding.

Commissioners of mental health services should ensure any service specifications take into account the recommendations in this guideline when it is finalised.

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## 1.1  First contact

These recommendations are for all services and providers.

1.1.1 Ensure all staff working in health (including urgent care), social care, voluntary and community sectors, and the criminal justice system[^1] who may come into contact with young people and adults with severe mental illness who misuse substances (dual diagnosis):

- understand that it is important to meet the needs of people with dual diagnosis wherever they present because they may have chaotic lifestyles and are often excluded from accessing services
- help people with any urgent physical health, social care, support or housing needs, either directly or by putting them in touch with other services.

1.1.2 Ensure the person is referred to secondary care mental health services for assessment and care planning (see section 1.2).

[^1]: NICE is developing a guideline on the mental health of adults in contact with the criminal justice system.
1.1.3 Be aware that people with dual diagnosis may have a range of physical health conditions including:

- cardiovascular, respiratory or hepatic conditions or related complications
- infectious disease
- cancer
- problems with blood glucose management.

1.1.4 Be aware that unmet social care needs such as social isolation or poor housing may lead people with dual diagnosis to have a relapse or affect their physical health.

1.2 Care planning

These recommendations are for all practitioners involved in developing and delivering a care plan.

On admission to secondary care mental health services

1.2.1 Develop a care plan after admission into secondary care mental health services (in line with the Care Programme Approach) for each young person or adult that:

- assesses the person’s social care, physical and mental health needs and any substance misuse problems they may have
- involves practitioners who can meet the person's needs from health and social care disciplines, for example: medicine, nursing, social work, occupational therapy and housing
- lists how the person's identified needs will be met and reviewed at every contact
- includes how the person can help implement the care plan.

Also see NICE's guideline on psychosis with coexisting substance misuse (recommendations 1.2.1–1.2.2 and 1.4.10–1.4.14).

1.2.2 Provide the person with a care coordinator within mental health services in the community (in line with the Care Programme Approach) to act as a
contact for the person, their family or carers and to help with developing and coordinating the care plan.

1.2.3 Ensure the care plan outlines how the safeguarding needs of all young people and adults with dual diagnosis will be met. (See also recommendations on safeguarding issues 1.1.14; and 1.1.16–1.1.20 in the NICE guideline on psychosis with co-existing substance misuse.)

1.2.4 Share the care plan with the person and other services as needed (see section 1.3 for recommendations on confidentiality and data sharing).

1.2.5 In line with the Care Act 2014 ensure carers who are supporting people with dual diagnosis are aware they are entitled to an assessment of their own needs. Ensure the assessment:

- recognises that carers have needs in their own right
- offers support for the effects of caring on the carers' mental health
- takes into account carers' views about services that could help them maintain their caring role and live the life they choose
- involves cross-checking any assumptions the person with dual diagnosis has made about the support their carer will provide
- advises carers that they may also be entitled to a personal budget to buy care to support them, for example to have a break from their caring responsibilities
- gives information and advice on how to access services in the community.

Involving people in care planning

1.2.6 When developing and reviewing the care plan:

- involve the person and their family or carers (where appropriate) to ensure the care plan is tailored to meet the person's needs (see also recommendation 8 in NICE’s guideline on behaviour change: individual approaches)
• discuss with the person how their abilities, strengths (such as the
  extent to which they can take part in activities of daily living) and their
  past experiences can help support their engagement and recovery
• take into account the concerns of the person’s family or carers
• recognise that the goals the person may have decided for themselves
  may differ from those identified by their service or provider
• jointly decide with the person which services would best meet their
  needs
• stay positive when talking about the prospects of recovery with the
  person, their family or carer.

See NICE’s guideline on psychosis and co-existing substance misuse and
service user experience in adult mental health for the principles of using a
person-centred approach.

1.2.7 Consider the following approaches to help young people and adults with
dual diagnosis remain involved in their care plan:

• one-to-one support
• training in self-care skills
• providing practical help with tasks that are important to the person
• providing help for example:
  – arranging or travelling with the person to hospital outpatient
    appointments or attendance at support groups
  – arranging for an advocate to accompany them at their appointments
    (see section 1.5 for recommendations on encouraging people to stay
    in contact with services).

Liaising with other organisations to meet physical health, social care, housing
or support needs

1.2.8 Adopt a coordinated approach (with shared responsibilities and regular
communication) when working with other organisations to meet people’s
wider health and social care needs.
1.2.9 Use every opportunity (including direct referrals if possible) to ensure people have prompt access to local services and actively follow them up to make sure their needs are being met.

1.2.10 As part of developing and reviewing the care plan, the person's care coordinator needs to:

- work with relevant primary care staff to meet the physical health needs of people with dual diagnosis; consider involving staff in substance misuse services
- work with relevant staff in local authorities, social care, community or voluntary sector organisations to meet the social care, housing or support needs. This can include:
  - personal care and hygiene
  - family and personal relationships
  - housing
  - employment
  - childcare responsibilities
- ensure an assessment of social care needs is carried out (in line with the Care Act 2014).

1.2.11 Consider covering the following health behaviours in the care plan:

- diet
- physical activity
- alcohol
- smoking
- other risky behaviours (see NICE's guideline on sexually transmitted infections, and the NICE pathways on hepatitis C and needle and syringe programmes).

1.2.12 Consider incorporating activities that can help to improve wellbeing and create a sense of belonging or purpose into a person's care plan. For example, encourage sport or recreation activities or attendance at
community groups that support good physical health. Ensure activities take account of a range of different needs.

Consider a range of approaches to encourage young people and adults with dual diagnosis to use services in the community. This could include, for example, the gym, education opportunities, volunteering or use of personal budgets (where applicable) for learning new skills such as return to employment.

Consider the suitability of the type of support that is available (for example, the type of housing, employment, detox or rehabilitation services) and take the person's preferences into account.

Recognise that people will have differing housing needs (for example high to low support or independent tenancies). Ensure safeguarding is in place.

Hold multidisciplinary case review meetings annually as set out in the Care Programme Approach or more frequently, for example every 3 months based on person's circumstances. Involve practitioners from a range of disciplines, including:

- secondary care mental health
- substance misuse
- primary care
- social care.

Check the person's physical health at least annually.

Consider reviewing the person within 3 months of them developing a new physical health problem (including monitoring for any adverse effects from medications).

Ensure the person's care plan is updated in response to changing needs or circumstances, including their social care, support or housing needs.
Discharge or transfer

1.2.20 Before the person is discharged or transferred between services (including from inpatient care to the community):

- invite any new practitioners who will be involved in the person's care to the multidisciplinary case review meetings and the discharge or transfer meeting.
- ensure the person's care plan includes strategies for ongoing risk management and details how they can get back in contact with services.

1.2.21 Ensure support is in place to help young people with dual diagnosis move to adult health or social care services. (See also NICE's guidelines on psychosis and coexisting substance misuse recommendations 1.8.1–1.8.9 and transition from children’s to adult services.)

1.3 Partnership working

These recommendations are for all services and providers.

1.3.1 Services in mental health, substance misuse, primary care, social care and support services need to collaborate with each other and other organisations in the community and voluntary sectors to provide a broad range of flexible services for young people and adults with dual diagnosis.

1.3.2 Services could consider working together to proactively encourage people with dual diagnosis to engage with services. This could be done:

- within an agreed set of local policies and procedures that is regularly reviewed by key strategic partners
- with good level of communication between all practitioners and a willingness to work across traditional institution boundaries
- by being responsive to requests for advice and joint-working arrangements
- by taking a shared response to risk management.

1.3.3 Ensure joint working arrangements are in place. This may include:
• services designed to ensure continuity of care and service provision
  (for example, when commissioning contracts are due to expire)
• services based on a local needs assessment
• ensuring the needs of young people and adults with dual diagnosis are
  part of other local needs assessment strategies, for example, on
  housing, employment projects, alcohol, drug services or crime
  prevention
• ensuring service quality is monitored and data sharing protocols are in
  place (see also recommendations 1.3.6–1.3.8).

1.3.4 Agree joint pathways to:

• meet the health, social care or other support needs and preferences of
  people with dual diagnosis, wherever they may present
• give people access to a range of primary health care providers and
  social care providers including GP practices, pharmacies, podiatrists
  dentists, social workers, housing or benefit advisers.

1.3.5 Ensure referral processes and care pathways within and across agencies
are consistent and that governance arrangements are in place. This
includes local care pathways to meet the physical health, social care,
housing and support needs of people with dual diagnosis.

Information sharing

1.3.6 Agree a protocol for information sharing between secondary care mental
health services and substance misuse, health, social care, education,
voluntary and community services (see the Caldicott Guardian Manual
2010).

1.3.7 Ensure that services have a consistent approach to getting people with
dual diagnosis help from the most relevant service by:

• sharing information on support services between agencies
• ensuring providers know about and can provide information on the
  services
1. taking up the responsibilities agreed in referral processes, providing timely feedback and communicating regularly about progress.

1.3.8 Ensure providers share information on how to manage challenging or risky situations (see also NICE's guideline on violence and aggression: short-term management in mental health, health and community settings).

1.4 Improving service delivery

These recommendations are for all services and providers responsible for delivery of services.

Adapting existing services

1.4.1 Adapt existing services rather than creating a specialist dual diagnosis service.

1.4.2 Make sure interventions that aim to improve the uptake of services, support harm reduction, change behaviour and prevent relapse are part of the service offered (see NICE's pathways on: psychosis and co-existing substance misuse; psychosis and schizophrenia in young people and adults; bipolar disorders; alcohol misuse and drug misuse).

1.4.3 Offer the person individual, face-to-face or phone appointment sessions to help encourage people with dual diagnosis to use services. Offer phone sessions to their family or carers. Sessions could cover:

- how the person is coping with their current mental health and substance use and its impact on their physical health and social care needs
- progress on current goals or changes to future goals
- ways to help the person stay safe
- monitoring symptoms
- getting support from family, carers or providers.

Determine how often the sessions take place based on the person's needs.
1.4.4 Consider the following:

- contingency plans within services to help the person (and their family or carers) with a potential crisis and ensure these are updated to reflect changing circumstances
- support to sustain change and prevent relapse
- discharge planning, including planning for potential relapses so that the person knows which service to contact and the service has the information needed to provide the right ongoing support. (See also NICE's guideline on transition between inpatient hospital settings and community or care home settings for adults with social care needs.)

Making services inclusive

1.4.5 Ensure existing services are adapted so that they can better engage and meet the needs of young people and adults with dual diagnosis.

1.4.6 Involve young people and adults with dual diagnosis, their family or carers in improving the design and delivery of existing services to provide a good standard of care (see section 1.2). This may include developing interventions and training, taking part in steering meetings and giving feedback on services.

1.4.7 Provide local services in places that are easily accessible, safe and discreet. Bear in mind any perceived stigma involved in being seen to use the service. Consider flexible opening times, drop-in sessions, or meeting people in their preferred locations.

1.4.8 Ensure people with dual diagnosis, their family or carers are given accurate information about relevant local services (for example, community groups or family support groups). This could include information on how to access services, ways to contact the service, opening hours and how long the waiting list may be.
Support for staff

1.4.9 Ensure the care coordinator in secondary care mental health services is supported to provide or coordinate flexible, personalised care based on a range of existing services (see section 1.5).

1.4.10 Recognise that different attitudes towards mental health and drug- or alcohol-related problems may exist between agencies and that this may present a barrier to delivering services. To overcome this:

- challenge negative attitudes or preconceptions about working with people with dual diagnosis
- develop leadership skills so staff can challenge attitudes and preconceptions (for example see dual diagnosis capability framework).

1.4.11 Ensure practitioners have the resilience and tolerance to help people with dual diagnosis through relapse or crisis so that they are not discharged before they are fully equipped to cope or consequently excluded from services.

1.5 Encouraging people to stay in contact with services

These recommendations are for all services and providers.

Building relationships

1.5.1 Recognise that building a relationship with people who have a dual diagnosis, may take time and involves:

- showing empathy and using a non-judgemental approach to listen, identify and be responsive to the person's needs and goals
- providing consistent services, for example, where possible keeping the same staff member as their point of contact and the same lead for organising care
- staying in contact by using the person's chosen method of communication (for example, by letter, phone, text, emails or outreach work, where possible).
1.5.2 Explore and discuss with the person if there are any reasons why they may be unwilling to use services to improve their physical health, or to receive social care support. This may include:

- the way services are organised
- inability to attend services, because they are not convenient (for example, services are not local, transport links are poor, or do not provide childcare)
- fear of stigma, prejudice or of being labelled as having both mental health and substance misuse problems
- feeling coerced into using treatments or services that do not reflect their preferences or their readiness to change
- previous poor relationships with practitioners
- other personal, cultural, social, environmental or economic reasons.

1.5.3 Help those with dual diagnosis who may find it difficult to engage with services to get into and stay connected with services by initiating and maintaining contact using proactive and flexible approaches (see recommendation 1.2.6).

1.5.4 Recognise that people with dual diagnosis are at higher risk of not using or losing contact with services, including for example:

- people who are homeless
- people who have experienced or witnessed abuse or violence
- people who are young
- men
- people who are parents or carers who may fear the consequences of contact with statutory services.

**Non-attendance**

1.5.5 Ensure non-attendance at an appointment is viewed by all practitioners as a matter of concern. Discuss and agree what follow-up actions should be taken with key practitioners in secondary care mental health services involved in the person's care plan. This could include:
• contacting the person's care coordinator within mental health services in the community immediately (particularly if there is a risk of self-harm or suicide) or at least within 24 hours, if there are existing concerns
• visiting the person at home, contacting any other practitioners involved in the person's care (identified in the person's care plan; see recommendation 1.2.1), or contacting family members
• ensuring the person is not automatically discharged without a discussion with the person's care coordinator and with all the practitioners involved in the person's care.

Putting this guideline into practice

NICE has produced tools and resources [link to tools and resources tab at publication] to help you put this guideline into practice.

Putting a guideline fully into practice can take time. How long may vary from guideline to guideline, and depends on how much change in practice or services is needed. Implementing change is most effective when aligned with local priorities.

Changes should be implemented as soon as possible, unless there is a good reason for not doing so (for example, if it would be better value for money if a package of recommendations were all implemented at once).

Different organisations may need different approaches to implementation, depending on their size and function. Sometimes individual practitioners may be able to respond to recommendations to improve their practice more quickly than large organisations.

Here are some pointers to help put NICE guidelines into practice:

1. **Raise awareness** through routine communication channels, such as email or newsletters, regular meetings, internal staff briefings and other communications with all relevant partner organisations. Identify things staff can include in their own practice straight away.

2. **Identify a lead** with an interest in the topic to champion the guideline and motivate others to support its use and make service changes, and to find out any significant issues locally.
3. **Carry out a baseline assessment** against the recommendations to find out whether there are gaps in current service provision.

4. **Think about what data you need to measure improvement** and plan how you will collect it. You may need to work with other health and social care organisations and specialist groups to compare current practice with the recommendations. This may also help identify local issues that will slow or prevent implementation.

5. **Develop an action plan** with the steps needed to put the guideline into practice, and make sure it is ready as soon as possible. Big, complex changes may take longer to implement, but some may be quick and easy to do. An action plan will help in both cases.

6. **For very big changes** include milestones and a business case, which will set out additional costs, savings and possible areas for disinvestment. A small project group could develop the action plan. The group might include the guideline champion, a senior organisational sponsor, staff involved in the associated services, finance and information professionals.

7. **Implement the action plan** with oversight from the lead and the project group. Big projects may also need project management support.

8. **Review and monitor** how well the guideline is being implemented through the project group. Share progress with those involved in making improvements, as well as relevant boards and local partners.

NICE provides a comprehensive programme of support and resources to maximise uptake and use of evidence and guidance. See our [into practice](#) pages for more information.

Also see Leng G, Moore V, Abraham S, editors (2014) Achieving high quality care – practical experience from NICE. Chichester: Wiley.

**Context**

Adults and young people who have a severe mental illness and misuse substances (defined as having a dual diagnosis in this guideline) have some of the worst health,
wellbeing and social outcomes (Relationship between dual diagnosis: substance misuse and dealing with mental health issues Social Care Institute for Excellence).

It is not clear how many people in the UK have a dual diagnosis because of several factors, including the fact that some people in this group do not use services or get relevant care.

The Department of Health’s Refocusing the Care Programme Approach identifies people with dual diagnosis as one of the groups in need of an enhanced Care Programme Approach because they are not being identified consistently and services are sometimes failing to provide the support they need. The policy highlights the need for a whole systems approach to their care, involving a range of services and organisations working together. This guideline aims to address this need.

Groups covered in this guideline include: young people (aged 14–25) and adults who have been diagnosed as having a severe mental illness and who misuse substances and who live in the community. The age cut-off for young people has been set at 14 to reflect the small numbers affected below this age – and the fact that many early intervention services usually start at age 14.

In this guideline, severe mental illness includes a clinical diagnosis of:

- schizophrenia, schizotypal and delusional disorders, or
- bipolar affective disorder, or
- severe depressive episode(s) with or without psychotic episodes.

Substance misuse refers to the use of legal or illicit drugs, including alcohol and medicine, in a way that causes mental or physical damage.

More information

To find out what NICE has said on topics related to this guideline, see our web pages on alcohol or drug misuse and mental health and behavioural conditions. For specific recommendations on monitoring and promoting recovery of physical health see our guidelines on psychosis and coexisting substance misuse, psychosis and schizophrenia in adults, psychosis and schizophrenia in children, bipolar disorder.
The committee’s discussion

Evidence statement numbers are given in square brackets. For an explanation of the evidence statement numbering, see the evidence reviews section.

Section 1.1 First contact

The discussion below explains how we made recommendations 1.1.1–1.1.4.

The committee noted that people with dual diagnosis are a vulnerable group, who often have poor physical health, are unemployed, homeless or are at risk of other people taking advantage of them. The committee was aware from the evidence (review 1) and their experience that people with dual diagnosis may present in different settings, so recommended that all staff coming into contact with them should be able to understand their needs and help them access services. The committee members were aware from their experience that people may present in crisis (for example at A&E) but they may be also be found opportunistically in other settings and identified as needing immediate assistance with a range of needs, including their mental or physical health, substance misuse or social care needs. They noted that the physical health and social care needs of people with dual diagnosis are often overlooked because of the challenging nature of dealing with both mental health and substance misuse issues. They also noted that people with dual diagnosis are often excluded from services because no one wants to take responsibility for them and that they need help to access a wide range of services.

The committee members were aware that criminal justice system settings were not included in the scope. However, they felt it was important to highlight this setting in recommendation 1.1.1 because it is a potential route for people with dual diagnosis to come into contact with healthcare services. This was also reflected in the expert testimony the committee heard on primary care services for homeless people (EP4).

The committee noted that wherever people with dual diagnosis present a similar approach to helping them access care is needed. The committee advised that secondary care mental health services needs to be the lead organisation responsible
for delivery of services and therefore made a recommendation to refer people with
dual diagnosis to secondary care mental health services.

The committee heard from an expert [EP4] on the physical health issues that can
affect people with dual diagnosis. They noted that although the expertise was from a
perspective of primary care services for homeless people, it was felt that the range of
health needs identified could be transferable to the wider dual diagnosis population.
The committee noted the evidence from 1 low quality [-] UK qualitative study which
highlighted commissioners’ views that the health and wellbeing of people with dual
diagnosis are not being met [ES2.1.2] and also reflected on the gap in the evidence
[ES1.1.8] on the prevalence of coexisting physical health problems. The committee
agreed to make a research recommendation in this area.

The committee noted that because of the complexity of their needs, people with dual
diagnosis are at increased risk of poor self-care, losing contact with family and
friends, social isolation or living in poor housing or having their homes abused by
others as venues for substance misuse or drug dealing.

From moderate to strong evidence from 4 cohort and 6 case-control studies, the
committee members were aware of the range of social care needs of people with
dual diagnosis in the UK [ES1.1.9]. They were also aware from expert testimony
[EP2] and their own experience of working with people with dual diagnosis of the
detrimental effects poor or unmet needs (such as social isolation or poor housing)
can have on person’s health and recovery process. This was supported by evidence
from 1 high quality and 1 moderate quality study [ES2.2.1]. They noted that unmet
needs could lead to relapse, using substances, deterioration in mental health,
offending behaviour or could also affect physical health.

**Section 1.2 Care planning**

The discussion below explains how we made recommendations 1.2.1–1.2.21.

**On admission to secondary care mental health services**

The committee agreed that secondary care mental health services need to take the
lead in coordinating services. The committee noted that care planning is usually led
by a care coordinator because this is part of the Care Programme Approach.
However, they noted that care coordinators are part of a multidisciplinary team subject to supervision and responsibility would lie with a consultant.

Based on their experience, the committee members advised that a care coordinator within mental health services in the community is assigned once a person has been referred into secondary care mental health services. The committee was also aware from evidence review 2 that having a continuity of contact with a key contact encourages people to keep in touch with services. Based on their expertise the committee made a recommendation that a care coordinator can take the key role in developing and reviewing a person’s care plan. The committee noted that the care coordinator would be responsible for organising delivery of range of needs including mental health, substance misuse, access to health or social care services with the support of a wider team and with supervision.

The committee noted from 1 moderate quality study and 1 low quality UK study [ES2.1.1] and their own experience that timely assessments can help people with dual diagnosis to access services and maintain engagement with their care plan. The committee agreed with the recommendations in NICE’s guideline on *psychosis with co-existing substance misuse* on the principles of recognition and assessment and felt it would be appropriate to link with this guideline. They recognised that the psychosis guideline has a narrower focus than this dual diagnosis guideline.

The committee members were aware from their practice and evidence [ES2.1.10] that services often fail to take responsibility for people with dual diagnosis. The effects of this failure can be far-reaching for example, causing the person to relapse, have physical health problems because of poor housing, or become unwilling to engage with services [ES2.2.1, ES2.2.2, ES2.2.4]. The committee was mindful that there could be cases where a person may have issues with both poor housing and physical health and that this may not always be a ‘cause–effect’ relationship.

The committee was aware from their experience of the importance of highlighting safeguarding issues for this vulnerable population. The committee was also aware of current legislation ([Care Act](https://www.gov.uk/care-act-2014) 2014) that entitled carers to have an assessment of their needs and they were aware from the evidence this may be particularly be the case if the carers are children [ES2.1.9; 1 UK study of low quality]. The committee therefore
developed a recommendation based on their expert knowledge to highlight young and adult carers' needs and ways to support carers for this specific population.

**Involving people in care planning**

The committee took into account qualitative evidence from 2 moderate quality studies and 1 low quality UK study [ES2.2.9] which highlighted that an approach that encouraged the person to be involved in their care plan decisions and respected their preferences can help the person adhere to the care plan. The committee was mindful that changes to behaviour may be a lengthy process and advised that NICE's [behaviour change: individual approaches](#) guideline may provide useful strategies.

The committee reflected on their experience and noted that providers need to understand what is having an effect on the person each time they see them so that they can provide the right level of support to the person at each time point. The committee reflected on their experience and noted that people can recover. The committee members also noted that in people with dual diagnosis the notion of recovery may not necessarily be about reducing their substance use but about leading a productive life. They were aware from the evidence [ES2.1.4] and their experience that the nature of relationships between a health or social care professional and a person with dual diagnosis can impact on a person’s willingness to engage, their response to care and approach to recovery. They felt that although recovery may take time, all interactions providers need to convey a sense of hope and optimism that it is possible.

The committee members were aware from their experience the importance of a person-centred approach. This was reinforced by the review on views and experiences of providers, commissioners, people with dual diagnosis, family and carers [review 2] and their experience. The committee was also aware of recommendation 1.1.1 in NICE’s guideline on [psychosis with co-existing substance misuse](#) which outlines adopting a non-judgemental and empathetic approach built on trust and respect. The committee felt it was important to take a person-centred approach when developing and reviewing the care plan [ES2.1.4, ES2.2.4, ES2.2.7] and recommended involving people in their care planning.
The committee recommended practical strategies that may help improve uptake of services and prevent relapse. This was based on evidence from 1 moderate quality study which noted self-care skills helped with daily living [ES2.2.3], expert testimony [EP2] and the committee's expertise.

**Liaising with other organisations to meet physical health, social care, housing or support needs**

The committee noted from the evidence [ES2.1.7] from 1 high, 2 moderate and 3 low quality qualitative studies (4 in the UK) that a lack of a shared approach could act as a barrier to providing health and social care services. The committee heard from an expert on local partnership working [EP1] and experts working with people with dual diagnosis who are homeless [EP2]. The experts highlighted factors that could help with a coordinated approach. Based on the evidence, the expert testimonies [EP1, EP2] and their own experience, the committee was in agreement that important factors in providing a coordinated approach included a shared vision, joint responsibilities and regular communication.

The committee members also highlighted the importance of prompt access in this recommendation based on their own experience and findings from qualitative evidence from 1 low quality UK study [ES2.2.8]. The committee members felt that direct referrals may be useful way to ensure timely response to the needs of this group. They noted that direct access to services may be beneficial (compared to for example, open access drop-in clinics) because this would give the person a sense of continuity of care which in turn may also enhance feelings of trust [ES2.2.4].

The committee developed recommendations to highlight the range of agencies or providers people in secondary care mental health services (for example, a care coordinator) would need to work with to ensure people with dual diagnosis receive care for their wider health or social care needs. The committee also highlighted in the recommendations physical health and social care needs that need to be taken in consideration as part of developing and reviewing a care plan.

The committee reflected on the expert testimony [EP4], their knowledge and experience to highlight how a person’s physical health could be improved and provided examples of how this may be achieved. This includes improving health
behaviours (such as diet, smoking or physical activity) and minimising risky
behaviours (such as unprotected sex, sharing needles). It was mindful this is not an
exhaustive list and recognised other behaviours may need to be addressed in the
care plan.

The committee also noted the importance of encouraging involvement in activities to
improve physical wellbeing (for example football activities or walking groups) but
were aware the risk of widening inequalities if activities only reach people who
already use services. The committee agreed that potential inequalities could be
addressed by recommending providing inclusive services and strategies to access
services to improve engagement with services.

In relation to social care, living or housing-related needs, the committee developed a
list of social care, housing and support needs (in line with the Care Act 2014).

The committee reflected on the evidence [ES1.1.9, ES2.2.1, ES2.2.2, ES3.1], expert
testimony [EP2] and their own experience to inform their recommendations on social
care, housing or support needs. The committee noted from the evidence [ES3.1] and
members' experience that a modest improvement in outcomes could be made in
areas such as in housing, employment or social functioning. In relation to housing
needs, the committee members noted there was strong evidence [ES1.1.9] from a
meta-analysis of 3 cohort and case-control UK studies (2 high quality; 1 low quality)
that people with dual diagnosis (compared with those with severe mental illness
only) had increased probability of a history of homelessness or housing problems.
There was also evidence from 1 high quality UK case-control study that people with
dual diagnosis (compared with severe mental illness only) had an increased
probability of living in the most deprived areas. The committee noted there was
moderate evidence from 3 high quality UK cohort studies that showed a greater
number of people with dual diagnosis are unemployed compared to those with
severe mental illness only. The committee also noted the evidence for social
functioning outcomes was mixed. A meta-analysis of 2 UK case-control studies (1
high and 1 moderate quality) showed no difference in social functioning between a
group with dual diagnosis and severe mental illness only. However, 1 high quality UK
cohort study showed poor social functioning in people with dual diagnosis compared
to those with substance misuse. It was noted the evidence was mainly from people
in contact with secondary care mental health services and may not adequately reflect the needs in the wider dual diagnosis population. They also were aware that unmet needs (such as housing) can also have a detrimental impact on a person’s health and recovery process [ES2.2.1].

Recommendations on approaches to encourage use of services and the suitability of the type of support were based on committee’s expertise and EP2. The committee was aware that people with dual diagnosis are particularly at risk of being taken advantage of and highlighted the importance of safeguarding in relation to housing needs.

**Review**

The committee members were aware from their experience and evidence from 1 high quality, 6 moderate quality, and 2 low quality UK qualitative studies of the benefits of an integrated approach to care [ES2.2.6]. They noted that this could increase engagement and result in positive change in health, functioning and wellbeing. The committee also took into account the evidence from 3 moderate quality qualitative studies and 1 recent UK study (of low quality) in a voluntary sector organisation [ES2.1.7], members' experience, and expert testimony [EP2]. The committee noted the importance of different disciplines working collaboratively, and taking part in case review meetings. Although the UK studies set in voluntary sector services (included in ES2.1.7 and ES2.2.6) were low quality, the committee felt the findings were relevant as they reflect views of providers in or users within this setting.

The committee noted that the frequency of case review meetings would vary and would involve multidisciplinary team members to make sure a person’s care plan is up to date and relevant. The recommendation to review annually was based on the Care Programme Approach, but the committee recognised that review of care plan is indicated by the person's level of need and circumstances.

The committee noted the importance of regular monitoring of physical health (based on EP4), including for adverse effects of medications. The committee was aware there was strong evidence from 3 UK case control and cohort studies [ES1.1.8] to show that people with dual diagnosis (compared with those with severe mental
illness-only) are less likely to adhere to medications. The committee heard in expert testimony [EP4] about side effects of medication and were aware from their experience that this includes effects such as weight gain or adverse effects. They felt this could be a barrier to adhering to treatment and could have a negative impact on a person’s mental or physical health. The committee’s recommendation on frequency of monitoring was based on their expertise.

The committee members acknowledged that the findings from the evidence on working collaboratively and the views expressed in the expert testimony [EP4] reflected their experiences of working with people with dual diagnosis. They noted that changes in circumstances need to be taken into account in a person’s care plan and physical health or social care, support or housing needs revised accordingly.

Discharge or transfer
The committee members noted that transfer or discharge are key points when a person can lose touch with services. They noted from the evidence [ES2.2.1] on challenges people can face when transitioning between services and felt this was applicable to other transitional contexts, for example, when people transition from inpatient care to community settings. Therefore they felt that handover of care upon discharge or when a person transfers to another service (in consultation with other providers) were important points to be included in the recommendation.

The committee heard from an expert in early intervention services [EP3] and noted the importance of making sure referral for transition to adult services for young people was identified in the guideline.

Section 1.3 Partnership working
The discussion below explains how we made recommendations 1.3.1–1.3.8.

The committee decided to recommend partnership working because provision of health and social care services for people with dual diagnosis is often fragmented and inconsistent. This was based on their experience and the evidence from 7 studies (2 high quality, 3 moderate quality and 2 UK studies of low quality) [ES2.2.4].

The committee advised that from April 2013 there have been separate funding streams for mental health and substance misuse services, with mental health
services funded by clinical commissioning groups and substance misuse services by local authorities. The committee felt this reflected a long standing division between the mental health and substance misuse and has led to 2 different sets of organising paradigms for commissioners which does not serve people with dual diagnosis. They also noted an additional challenge for addiction services is that although funding is from local authority budgets they are subject to commissioning and competitive tendering.

The committee reflected on current practice. The committee noted that although the Department of Health’s 2002 policy guide had set the vision for how services and care for people with dual diagnosis could be delivered, implementation was lacking. The committee was aware of Public Health England guidance on Co-existing alcohol and drug misuse with mental health issues: guidance to support local commissioning and delivery of care [to be published June 2016] which has set out the importance of joint working.

The committee noted from the qualitative evidence including 3 UK studies (1 high and 2 low quality studies) in a range of settings [ES2.1.7] that different disciplines working together, with a shared approach in their dealings with people with dual diagnosis and shared responsibilities, could facilitate the delivery and improve the quality of health and social care services. This could be done by using a shared approach, co-management of cases, and regular communication. The committee also heard from an expert in local partnership working [EP1] who described a framework designed to help local areas design and deliver flexible and coordinated services for people experiencing multiple needs.

The committee noted that there needs to be a strategic framework for dual diagnosis work to operate within, with buy-in from providers and commissioners, for services to collaborate locally. Based on the evidence, the expert testimony [EP1] and their own experience, the committee members were in agreement that a cross-sector partnership, with a shared understanding of the problem (based on assessment of local needs), and a shared vision for the future were important factors. Based on their expertise and expert testimony, they developed a recommendation on how services need to work together. They noted the lack of evidence from review 1.2 on existing care pathways and felt this is an area for research recommendation.
The committee was aware of evidence from review 2 that a lack of policy on referrals has an effect on the organisation and continuity of care. Evidence from 4 qualitative studies conducted in different settings (including 1 UK study set in voluntary sector) noted uncertainty on who should make referrals can also have an impact [ES2.1.11]. The committee members noted that the evidence from qualitative studies [ES2.1.10, ES2.1.11] was consistent with their experience. The evidence noted that pathways were deemed to be inadequately planned and supported and that movement across a care pathway was often restricted because of failure in services to take responsibility. They also noted that continuity of care can be interrupted because of changes in the commissioning process or cycle. For example, they noted retendering for services can lead to disruption and the need to build new care pathways.

One UK low quality qualitative study exploring views of commissioners [ES2.1.11] noted that good links between statutory and voluntary sector improved outcomes, such as reduced waiting times and delivery of care. This could also help with organisation and continuity of care. The committee was also aware of evidence from the study that highlighted that existing resources were stretched [ES2.1.5] and that lack of funding was affecting provision in non-voluntary sectors [ES2.1.6]. The committee noted that this study was published in 2002. It noted service provision for addiction services has changed considerably in the past 10 years, the demography of the individuals, the treatment and the types of substances used will all have changed markedly since 2002.

The committee reflected on their experience and acknowledged the importance of including the needs of people with dual diagnosis in the joint strategic needs assessment and in a local strategy (for example, housing, alcohol or drug services, or crime prevention). The committee noted that referral processes and pathways need to be in place to ensure this happens. This would help with a joined up approach to meet the needs of people with dual diagnosis who often fall between the gaps in services. The committee noted from the evidence that currently there isn't a service configuration that is in place nationally (review question 1.2).

**Information sharing**

The committee made recommendations to highlight the importance of information sharing. The committee noted an expert testimony [EP2] highlighted that a barrier
the voluntary sector often faces is that confidentially is often used as an excuse not to share information. The committee also noted the importance of services having knowledge of other local services and being able to tell people with dual diagnosis about them [ES2.2.3, ES2.2.10]. For example, one UK low-quality study set in the voluntary sector noted GPs were unaware of local community groups which people with dual diagnosis could access [ES2.2.3]. The committee was also aware from its experience that as part of providing inclusive services staff need to be equipped with skills. For example, skills to deal with challenging nature of working with people who may be intoxicated.

Section 1.4 Improving service delivery

The discussion below explains how we made recommendations 1.4.1–1.4.11.

The committee decided to recommend improving delivery of existing services that are inclusive because the needs of people with dual diagnosis are often not taken into account and they face the risk of being excluded from mainstream services.

Adapting existing services

The committee agreed that the recommendations needed to focus on improving existing services instead of creating a specialist dual diagnosis service. They felt that the standard care delivered in the UK could be improved by increasing engagement with existing services and that existing capacity and resources could be used to deliver these in the system. The committee adopted this approach for the design of a service delivery model and developed the content of the service model based on the evidence, economic model (see economic considerations), expert testimony and their expertise.

The committee considered the evidence for the effectiveness and efficiency of service delivery models [ES3.1, ES3.2, ES3.3, ES3.4, ES3.5, ES3.6, ES3.7, ES3.8, ES3.9, ES3.10], which included randomised controlled trials and observational studies. The evidence covered a range of service delivery interventions, showing some positive outcomes and there was value in what the models were aiming to achieve. However, the members agreed that there was no overwhelming evidence of benefit to indicate a particular model should be recommended (please see below for committee’s interpretation of the evidence on the effectiveness, cost effectiveness
and efficiency of service delivery models). The committee members were also aware of moderate evidence from 12 UK studies [ES1.2.1] that there are inconsistencies in the configuration of dual diagnosis services within NHS trusts across the UK. These inconsistencies lie in a number of areas including sources of funding, structure of services, type of staff members, services delivered and coordination of care. The committee also considered the evidence on current configuration of services [ES1.2.1] and observed there were few specialist services for adults. They were also aware of evidence from 1 low quality UK study that described mixed views amongst staff in a specialist dual diagnosis service on whether dual diagnosis services should be separate or integrated with mental health or substance misuse service [ES2.1.13]. The committee also noted that there was evidence from the same study indicating that most commissioners felt that integrating services would be essential for the effective and efficient delivery of care for people with complex needs. Additionally, some commissioners noted that relationships between different services could be expected to improve if they were required to share budgets and resources.

The committee felt this finding should be treated with caution because the funding landscape has changed considerably since 2002 and their experience suggests that a third tier of provision may not necessarily meet the needs of people with dual diagnosis. Based on their experience, the committee members noted that 'integration' should be about joint working and coordinated care and did not feel a specialist service was the way forward for this group.

The committee also noted that there was limited description of the comparator arms (often described as ‘treatment as usual’) in the studies included in review 3 and that most of the studies were conducted in the US. The committee’s view was that ‘usual care’ in the US is likely to differ from that provided in the UK and the level of ‘usual care’ delivered in the UK was considered to be of a better standard. The committee’s expert knowledge and the evidence was used to develop a recommendation on aspects that could be included in a service. This includes interventions that have shown to be effective in NICE guidelines for either severe mental illness or substance misuse. The committee was aware of the Wenze (2015)[2] study included

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in the economic model. Members reflected on the components of the 'treatment-engagement' sessions in the Wenze (2015) study as well as their own experience to develop a recommendation on ways to improve engagement. The committee noted that any recommendation on improving service delivery needs to take into account the needs of those who reach crisis and those who relapse after discharge. This recommendation was based on their expertise. The committee was aware from the evidence and their experience that people experience fragmented care [ES2.2.4] and that plans need to be in place to allow people to move back into the system. It was noted that the Department of Health’s Mental Health Crisis Care Concordat would have information on developing an action plan to deal with people in a crisis.

The committee heard from an expert on the service delivery model in early intervention services [EP3]. The committee noted that these services are able to offer consistent and coordinated service because the staff working in these services have lower caseloads and so are able have more contact with the people they work with and provide stability.

**Making services inclusive**

The committee members were aware from their own experience, the evidence [review 1] and expert testimonies [EP1, EP2] of the benefits of supporting people with dual diagnosis to participate in improving services. The committee also noted from the evidence [ES2.2.9, ES2.2.10] the importance of involving people with dual diagnosis, and their family or carers and providing them with the information and support. The ways in which people with dual diagnosis, their family or carers could be involved in design and delivery of services was based on the findings from the review on epidemiology and current configuration [review 1].

The committee also observed that there appears to be an inequity in the way that people with dual diagnosis are treated by services compared with other groups. The committee’s experience and the evidence from the review of views and experiences of providers and commissioners and people with dual diagnosis [ES2.1.3, ES2.2.5] showed that this may be because of negative attitudes or stereotyping. A pessimistic attitude on the likelihood of the person staying in the service was also considered to be a contributor to this inequity.
The committee was aware from the evidence review on epidemiology that the prevalence of dual diagnosis varied across regions. Semi-rural areas seem to have the highest need yet specialist services are mostly in urban areas [ES1.1.2] and expert testimony showed high incidence of early psychosis in rural areas [EP3]. The committee agreed to not make a recommendation specifying content or configuration of service delivery by geographical settings, instead they felt that the most important message was to ensure that any services needed are delivered locally.

Although evidence on co-location of services (for example, services based in the same facility) was mixed [ES2.1.12], the committee members took into account the role of stigma in accessing services. The committee members were aware from their experience and from expert testimony [EP1; EP2] that people with dual diagnosis may be at risk of exploitation (for example, forced to become sex workers) or may have experienced trauma (for example, women may have experienced rape). The committee felt that was important that a recommendation was included in the guideline to highlight the importance of locating services in places that are safe and where there is minimal stigma attached to attending. The committee also considered the services highlighted in the evidence review (review question 1.2) on current configuration of services and developed a recommendation highlighting the importance of safety of location, low stigma and flexibility in opening times as factors that can help make services more accessible. The committee members were also aware from the evidence [review 2] that where people knew about services, barriers to access included difficulty in accessing services outside hours, long waiting lists and service not being local.

**Support for staff**

The committee noted the importance of support and supervision from their experience and the evidence from 2 high quality, 1 moderate quality and 2 low quality qualitative studies [ES2.1.15; 3 set in the UK]. Because of the complexity of the care coordinator's role the committee felt it was important to highlight in the recommendation the importance of a support structure for this role. The committee members were also aware from the evidence and their experience that addressing gaps in practitioners' knowledge can help with establishing links with other services and delivery of services [ES2.1.16].
Evidence from 1 high, 3 moderate, and 1 low quality qualitative studies [ES2.1.14; 2 set in the UK] found consistent views amongst providers and commissioners in a variety of settings that a barrier to service delivery or partnerships between services is different perceptions of drug and alcohol problems depending on the focus of the service. Services failing to take responsibility for people with dual diagnosis and the potential impact of this on meeting wider health, social care or support needs were highlighted in providers’ views across 6 qualitative studies [ES2.1.10]. Of these, 3 of the studies were set in the UK, 1 was of moderate and 2 were low quality. The committee noted although 1 of the UK studies was of low quality it was a recently published study and reflected the voluntary sector providers’ views. The committee drew on the evidence and their own expertise and noted that helping overcome negative attitudes in staff will help make sure people with dual diagnosis are not excluded from services.

The committee members were aware from the evidence [ES2.1.4] on the importance of establishing good relationships between practitioners and people with dual diagnosis. They noted there was high quality UK evidence from 1 study to show that practitioners perceived that behaviours such as misusing drugs could impact on relationships and act as a barrier to delivery of care. The committee reflected on the evidence and their experience and therefore recommended that there needs to be tolerance and resilience in services to work with people through relapse, poor attendance or a crisis as these can often lead the person being inappropriately discharged.

Committee’s interpretation of the evidence on effectiveness, cost effectiveness and efficiency of service delivery models (reviews 3 and 4):

The committee agreed that there was limited evidence of effect for assertive community treatment and integrated treatment interventions [ES3.1, ES3.2; ES3.3] in relation to mental health and substance misuse outcomes. The committee noted that fidelity to delivery of interventions (whether the intervention was delivered as designed) in the service models was reported for only 5 studies and where reported the fidelity was considered to be good. There was weak evidence for assertive community treatment based on 5 US RCTs, moderate evidence from 6 RCTs and 1 observational study (3 studies based in the UK) for integrated treatment interventions.
compared to treatment as usual. There was weak evidence from 1 RCT for
integrated treatment intervention compared to enhanced assessment and
monitoring. The RCTs did not all show a clear evidence of benefit. There was some
improvement noted in service use outcomes but members noted that it was
debatable whether this was necessarily an evidence of benefit. There was some
evidence of effect on social care outcomes such as housing, employment and social
functioning. The committee noted that the assertive community treatment
intervention model was no longer used in the US. The committee felt that although
the follow-up in these studies ranged from 24 weeks to 3 years, the length of time
needed to observe small improvements can sometimes take 5 to 10 years.

The committee considered the other service delivery interventions identified in
evidence review 3 (9 studies) including case management, contingency
management, staff training, or supportive housing [ES3.4, ES3.5, ES3.6, ES3.7,
ES3.8, ES3.9, ES3.10]. They noted that there was mainly weak evidence from small
studies, with short follow-up (ranging from 16 weeks to 78 weeks) and with the
exception of 3 studies (based in UK and Ireland) the remaining were mainly from US.
It was noted that fidelity to delivery of the intervention was reported in only 2 studies
(reported as low and high fidelity). The members discussed the potential value of
service models incorporating contingency management, peer support (delivered as
part of a care coordination intervention in 1 US study) or text messaging [ES3.5,
ES3.6, ES3.10] and considered these further under research recommendations.

The committee agreed that there was weak evidence for a staff training intervention
considered in the review of effectiveness of service delivery models [ES3.8] and
agreed not make a recommendation on this area. The committee noted that the 2
UK RCTs were of low quality, the evidence was inconsistent and did not appear to
show an overall benefit. In addition, a committee member reflected on their own
involvement in delivery of the intervention in one of the studies. It was noted there
were challenges in the delivery of a staff training intervention, particularly because of
staff moving between services, high turnover of staff and low fidelity to delivery of the
intervention.

The committee agreed there were several gaps in the evidence from review 3
including:
population (limited evidence on young people and vulnerable groups)

- interventions or measures (for example measures looking at improving accessibility and availability of services)
- outcomes (no evidence on physical health outcomes)
- efficiency of service delivery models - for example outcomes on accessibility of services (waiting times).

The committee discussed the evidence from the cost-effectiveness studies ES4.1, ES4.2, ES4.3, ES4.4, ES4.5 and the economic modelling [ES4.6] (see Economic considerations) when developing the recommendation on providing services.

**Section 1.5 Encouraging people to stay in contact with services**

The discussion below outlines how we made recommendations 1.5.1–1.5.5.

The committee members decided to make recommendations on encouraging people to stay in contact with services and making services accessible because they were aware from the evidence and their own experience that people with dual diagnosis may find it hard to initiate or maintain contact with services. Also, their physical health, social care, housing or support needs are not being met.

**Building relationships**

The committee noted from their experience that it is important to take a long-term and realistic view when working with people with dual diagnosis in relation to involving the person in their care plan and coordinating their care, particularly in light of the challenging nature of working with this group [ES2.1.8]. The committee members were aware from the evidence, their own experience, and expert testimony of the importance of continuity and flexibility in approaches to help people. The committee heard from experts working with people who are homeless [EP2] and reflected on their experience of range of methods that could be used to engage and stay in touch with a person. The committee considered the evidence from 4 qualitative studies (1 high; 1 moderate; 2 low quality), of which 2 were UK studies [ES2.2.4]. The evidence highlighted that a lack of continuity of care along with changes in staff can result in a lack of trust or an unwillingness to engage with services and good after-care was an important aspect of preventing relapse.
The members reflected on their experience and the evidence from 8 qualitative studies of mixed quality (2 high quality, 3 moderate and 3 low quality), with 3 of the studies (low quality) set in the UK. The studies examined the importance of relationships with healthcare professionals [ES2.2.7] and the committee agreed that a non-judgemental empathetic approach was needed when encouraging a person with dual diagnosis to stay in contact.

The committee noted that the evidence from views and experiences of people with dual diagnosis, their family or carers highlighted barriers to access or uptake of social care or physical health services [ES2.1.3, ES2.2.1, ES2.2.2, ES2.2.5, ES2.1.12]. Barriers included lack of support during a transitional period (for those who have previously had criminal convictions), failure to recognise cultural differences and mistrust of healthcare professionals or poor links to services. Negative connotations of being labelled as having problems with both mental health and substance misuse [ES2.2.5], negative attitudes, stereotyping or stigma about mental health diagnoses in substance misuse settings or about substance misuse in mental health settings [ES2.1.3, ES2.2.5] were also considered to be barriers to accessing services or receiving care.

The committee was aware from evidence review 2 and their experience that having continuity of contact encourages people to keep in touch with services. The committee recommended range of approaches based on their experience and expertise [EP2; EP4].

The committee recognised that all people with dual diagnosis face difficulties in receiving care [ES2.1.10] but wanted to highlight that some groups are particularly vulnerable. The committee noted there was moderate to strong evidence [ES1.1.5] that dual diagnosis is associated with those who were of younger age and male gender. The committee was mindful that homelessness is a frequent outcome for people with dual diagnosis. Factors contributing to this include people with dual diagnosis not being able get, or stay in contact with, the services they need. The committee reflected on their experience and also noted that a trauma history is almost always present in people with dual diagnosis, which can lead to disruptive attachments and associated challenging behaviour. The committee extensively discussed other groups but noted there were limitations in the evidence. They noted
that the evidence linking ethnicity with dual diagnosis was inconsistent [ES1.1.5].

The committee noted that apart from age, gender and ethnicity, there was a lack of evidence for particular groups identified in the equity impact assessment who are more likely to have a dual diagnosis. This includes, for example, people with a learning disability, teenage parents, Gypsies and Travellers, asylum seekers or refugees, lesbian, gay, bisexual, transsexual or transgender people, and sex workers [ES1.1.5]. The committee was aware from their experience that people with dual diagnosis have a range of social care needs, but noted that the evidence did not identify social care needs for particular groups identified in the equity impact assessment. For example, those who are socially isolated, on low income, have a history of being ‘looked after’ or are adopted or have a history of experiencing or witnessing domestic violence and abuse [ES1.1.9].

Although there was no evidence identified, the committee was aware from their experience that other groups may be reluctant to engage with or may encounter difficulties when engaging with services for people with dual diagnosis. These groups include for example people who are recent migrants, have language difficulties or are from specific religious communities.

Although it is not an exhaustive list, the committee highlighted the groups identified in recommendation 1.5.4 based on the evidence [ES1.1.5; review 2], their expertise and expert testimony [EP2].

The committee reflected on the evidence from review 2 and noted although the review provided insight to barriers and facilitators to delivery of care they felt there was research needed to understand the experience of people at different stages of recovery.

**Non attendance**

The committee members were aware from the evidence [ES2.2.7] and their experience that lack of emotional support and empathy can be a contributing factor to non-attendance. The committee were aware from their experience and the evidence [review 2] that non-attendance can often lead to discharge. Based on their expertise, they developed a recommendation that highlighted actions services can
take in collaboration with secondary care mental health services to ensure non-
attendance is treated as a matter of concern.

The committee reflected on their experience and expert testimony [EP2] and noted
the importance of maintaining contact and reaching out to people to help them
remain engaged with services. They reflected on their experience to suggest follow-
up actions in the recommendations to address non-attendance.

**Economic considerations**

An economic analysis was undertaken which comprised a review of existing cost
effectiveness studies and a bespoke economic model.

The findings from the review of evidence (from 1 UK and 7 US studies) were
inconsistent [ES4.1, ES4.2, ES4.3, ES4.4, ES4.5]. The US studies found that
integrated treatment leads to minor cost savings but the UK study found that the
intervention resulted in an increase in public sector costs. In all studies integrated
treatment appears to result in improvement in some outcomes; however economic
analyses used different outcome measures reported as changes on various scales
making comparisons challenging. Three studies adopted before-after design, studies
used different perspectives and time horizons, only 1 included economic study was
judged to be directly applicable, 3 studies were judged to be characterised by minor
limitations [++] , 4 studies by potentially serious limitations [+], and 1 study by very
serious limitations [−]. Overall, there is little evidence to support one service delivery
model over another, based on existing economic evidence.

The model was based on 3 studies. The first study, conducted in the US, comprised
a treatment engagement intervention (using resources more intensively compared
with standard care) for people with bipolar disorder and substance misuse. It was a
small study whose health outcome was inconclusive, but yielded resource use data.
The remaining 2 studies, both from the UK, were used to estimate baseline
admissions rates for people with dual diagnosis.

The model’s time-horizon was 1 year only. Consequently, increases in life
expectancy that might have occurred as a result of an intervention were not included
as benefits in the model. Because of the lack of data a further conservative 
assumption was that wider costs, particularly those falling on the criminal justice 
system, were not included. Further, the model's measured outcome might not have 
measured all of the health outcome benefits.

The model showed that an intervention which combined enhanced engagement with 
standard care would need to reduce relapses by about 12% for the intervention to 
become cost saving. However, the committee members had different views about 
whether UK standard care is better than that reported in the US studies. Assuming 
standard care in the UK is equivalent to the enhanced intervention modelled, it would 
be offering better outcomes at the same cost. By definition, that would be a cost 
effective approach. However, assuming standard care in the UK would need to be 
enhanced and therefore need additional resources, at a cost of £226 per individual 
and assuming an effect size of 10% the intervention would need to result in a small 
QALY gain of 0.002 (equivalent to 0.73 days in full health) for an ICER to be below 
the lower NICE cost-effectiveness threshold of £20,000 [ES4.6].

Given the results that were obtained even though a number of potential benefits 
were not considered because of the lack of data (for example on a person’s life 
expectancy, improvement in the substance misuse problem, improvement in the 
mental health of service users the reduction in health and social care and the 
criminal justice system costs) the treatment engagement intervention is very likely to 
be a cost-effective option.

Other points the committee discussed

The committee discussed the exclusion criteria in the scope and noted that exclusion 
of mental health disorders such as eating disorders was a major gap. The committee 
noted the criminal justice system settings were excluded from the scope but were 
aware of NICE guidelines currently in development that included this setting (mental 
health of adults in contact with criminal justice system and physical health of people 
in prison).

The committee considered a range of expertise that would be helpful to inform the 
development of the guideline and invited expert testimony in early intervention 
services, primary care, homeless, and local partnership working. The committee also
acknowledged other groups (refugees, veterans) but recognised that there were
general set of needs which would subsume the specific needs of particular
populations. The committee recognised criminal justice system settings were out of
scope but noted that young people and adults with dual diagnosis who need a safe
place to stay may come into contact with people within this setting, for example the
police. However the committee noted that resources for helping the police to support
people with vulnerabilities are available at Crisis Care Concordant.

The committee considered all the evidence available in developing this guideline.
However some evidence statements provided background information and could not
be explicitly linked to recommendations [ES1.1.1, ES1.1.3, ES1.1.4, ES1.1.6,
ES1.1.7]. The committee heard from an expert in early intervention services [EP5]
who described a study on contingency management (CIRCLE) which provided
background information and was not linked to a specific recommendation.

The committee discussed the various forms of support groups or mechanisms for
peer support. It was aware of mutual aid organisations including Alcoholics
Anonymous (AA), Narcotics Anonymous (NA) and SMART recovery and discussed
the merit of adding a reference to such forms of support as examples in the guideline
recommendations. It was also aware of the Public Health England guidance (A
briefing on the evidence-based drug and alcohol treatment guidance
recommendations on mutual aid) but noted it was not aware of evidence establishing
use of mutual aid in people with dual diagnosis. In addition because peer support
was an area identified for research recommendation, the committee did not
recommend specifying examples of mutual aid groups within the guideline
recommendations.

The committee also noted that there may be a stigma attached to the term
substance ‘misuse’ but recognised that this term is used in other NICE guidelines.

Evidence reviews
Details of the evidence discussed are in evidence reviews, reports and papers from
experts in the area.
The evidence statements are short summaries of evidence. Each statement has a short code indicating which document the evidence has come from.

Evidence statement (ES) number 1.1.1 indicates that the linked statement is numbered 1 in review question 1.1 of review 1. ES1.2.1 indicates that the linked statement is numbered 1 in review question 1.2 of review 1. ES2.1.1 indicates that the linked statement is numbered 1 in review question 2.1 of review 2. ES2.2.1 indicates that the linked statement is numbered 1 in review question 2.2 of review 2. ES3.1 indicates the linked statement is numbered 1 in review 3 and ES4.1 indicates the linked statement is numbered 1 in review 4. EP1 indicates that expert paper 1: ‘Local partnership working: examples drawn from the work of the Making Every Adult Matter coalition’ is linked to a recommendation. EP2 indicates that expert paper 2: ‘St Mungo’s: people who have a dual diagnosis and are homeless’ is linked to a recommendation. EP3 indicates that expert paper 3: ‘Early Intervention in Psychosis services’ is linked to a recommendation. EP4 indicates that expert paper 4: ‘Dual Diagnosis among homeless people: primary care perspective’ is linked to a recommendation.

If a recommendation is not directly taken from the evidence statements, but is inferred from the evidence, this is indicated by IDE (inference derived from the evidence).

Section 1.1: ES1.1.8, ES1.1.9, ES2.1.2, ES2.2.1; EP2, EP4; IDE

Section 1.2: ES1.1.8, ES1.1.9, ES2.1.1, ES2.1.4, ES2.1.7, ES2.1.9, ES2.1.10, ES2.2.1, ES2.2.2, ES2.2.3, ES2.2.4, ES2.2.6, ES2.2.7, ES2.2.8, ES2.2.9, ES3.1; EP1, EP2, EP3, EP4; IDE

Section 1.3: ES2.1.5, ES2.1.6, ES2.1.7, ES2.1.10, ES2.1.11, ES2.2.3, ES2.2.4, ES2.2.10; EP1, EP2; IDE

Section 1.5: ES1.1.5, ES1.1.9, ES2.1.3, ES2.1.8, ES2.1.10, ES2.1.12, ES2.2.1, ES2.2.2, ES2.2.4, ES2.2.5, ES2.2.7; EP2, EP4; IDE

Gaps in the evidence

The committee's assessment of the evidence on dual diagnosis identified a number of gaps. These gaps are set out below.

1. Evidence on the characteristics of people with dual diagnosis in the groups identified in the equity impact assessment. This includes for example, people with a learning disability, teenage parents, Gypsies and Travellers, asylum seekers or refugees, lesbian, gay, bisexual, transsexual or transgender people, and sex workers.

(Source review 1)

2. Social care needs of people identified in the equity impact assessment. This includes for example, those who are socially isolated, on low income, have a history of being ‘looked after’ or are adopted or have a history of experiencing or witnessing domestic violence and abuse.

(Source review 1)

3. Views and experiences of:
   a) commissioners
   b) primary care practitioners who work with vulnerable groups
   c) groups identified in the equity impact assessment (with the exception of young people and ex-offenders).

(Source review 2)

4. Interventions or measures assessing efficiency of services (for example measures looking at improving accessibility and availability of services).

(Source review 3)
5. Different models of service delivery (for example, a comparison of specialist, integrated or separate services) and efficiency of service delivery models.

(Source review 3)

**Recommendations for research**

The guideline committee has made the following recommendations for research.

1 **Epidemiology**

In the UK, how prevalent is a dual diagnosis and what are the physical health and social care, housing or support needs of people with this diagnosis?

**Why this is important**

There is limited evidence on the physical health and social care needs of people with dual diagnosis. This includes prevalence of coexisting physical conditions such as cardiovascular, respiratory or infectious diseases and social care needs such as social isolation or poor housing. Longitudinal evidence is needed to understand these. This will help design coordinated evidence based services to meet the wider health and social care needs of people with dual diagnosis and provide a good standard of care. People with dual diagnosis may present in a variety of settings and research that focuses on specific settings (for example, primary care) would be beneficial.

2 **What works?**

In the UK, what is the effectiveness and cost effectiveness of service delivery interventions (peer support, contingency management or text messaging delivered in conjunction with standard care) compared with standard care alone for young people and adults with dual diagnosis?

**Why this is important**

There is limited evidence on the optimal service delivery model to meet the needs of young people and adults with dual diagnosis. Evidence suggests there is increasing use of contingency management, peer support or text messaging as part of a service delivery intervention to help young people and adults with dual diagnosis access
services. Further research is therefore needed to assess their use, benefit and whether these have a role in improving engagement with services for people with dual diagnosis. Research is also needed on what a good outcome for this group is, the methodological challenges of encouraging research in people with dual diagnosis, and appropriate methodology for engaging with this group.

3 Evidence of cost effectiveness

What is the cost effectiveness of services (or elements of standard care) for young people and adults with dual diagnosis?

Why this is important

There is limited evidence on the cost effectiveness of interventions and services with this group. Further research is needed to investigate this and ascertain whether any particular services or elements of standard care provide better value for money.

4 Barriers and facilitators

What are the barriers and facilitators for young people and adults with dual diagnosis to obtain an optimal service (including optimal time frame for delivering interventions) to meet their needs and enable their recovery?

Why this is important

There is limited evidence that identifies the triggers for deterioration and the turning points for recovery for people with dual diagnosis. Although review 2 contains evidence on the views and experiences of people with dual diagnosis, their family or carers, it is not always clear to which point in the care pathway the views and experiences expressed relate to. As such, it is difficult to fully break down the experience of care received at various intervals along the care pathway. Understanding the experience of people who are at different stages of recovery and how they have maintained their progress and success (1 year, 3 years, 5 years, 10 years+) will help with designing more effective services and planning services that deliver interventions at the right time.
5 Care pathway

In the UK, what is the optimal care pathway for young people and adults with dual diagnosis?

Why this is important

There is a lack of published evidence on care pathways on treatment, management and follow up of people with dual diagnosis. In the UK, service configurations, treatment philosophies and funding streams act as barriers to coordinated provision of care. Separate mental health and substance misuse services that are often provided by different organisations, have different organisational and managerial structures, and staff within each service may often lack the knowledge and skills for working with people from another organisation. A review of what has worked or not in areas that have implemented changes to practice will help services develop optimal care pathways.

Glossary

Contingency management

Contingency management is a set of techniques that focus on changing specific behaviours. For example, in drug misuse, it involves offering incentives for positive behaviours such as abstinence or a reduction in illicit drug use, and participation in health-promoting interventions.