Safeguarding People Living with Dementia: How Social Workers Can Use Supported Decision-Making Strategies to Support the Human Rights of Individuals during Adult Safeguarding Enquiries

Jeremy Dixon 1*, Sarah Donnelly2, Jim Campbell2 and Judy Laing3

1Department of Social & Policy Sciences, University of Bath, Bath BA2 7AY, UK
2School of Social Policy, Social Work and Social Justice, University College Dublin, Dublin, Ireland
3University of Bristol Law School, Bristol, UK

*Correspondence to Jeremy Dixon, Department of Social & Policy Sciences, University of Bath, Bath BA2 7AY, UK. E-mail: j.dixon2@bath.ac.uk

Abstract

Dementia may make adults more susceptible to abuse and neglect and such mistreatment is recognised as a human rights violation. This article focuses on how the rights of people living with dementia might be protected through the use of supported decision-making within safeguarding work. The article begins by reviewing the aims and scope of adult safeguarding services. It then describes how the concept of ‘legal capacity’ is set out in the UN Convention on the Rights of Persons with Disabilities (CRPD) and how this differs from the concept of ‘mental capacity’ in the Mental Capacity Act 2005. Focussing on practice in England, it is argued that tensions between the CRPD and domestic law exist, but these can be brought into closer alignment by finding ways to maximise supported decision-making within existing legal and policy frameworks. The article concludes with suggested practice strategies which involve: (i) providing clear and accessible information about safeguarding; (ii) thinking about the location of safeguarding meetings; (iii) building relationships with people living with dementia; (iv) using flexible timescales; (v) tailoring information to meet the needs of people living with dementia and (vi) respecting the person’s will and preferences in emergency situations.
Introduction

Dementia is an umbrella term used to refer to a range of conditions leading to impairments in memory, language and sensory awareness. Whilst the causes of abuse and neglect are complex, research shows that older adults with dementia experience higher rates than those without dementia (Fang and Yan, 2018). Such mistreatment is recognised as a human rights violation by the World Health Organisation (WHO, 2016). Speaking at the Global Action Against Dementia conference in 2015 the UN Independent Expert on the Enjoyment of all Human Rights by Older People stated that:

the rights and needs of person’s with dementia have been given low priority in the national and global agenda. In particular, with the progression of the disease, as their autonomy decreases, persons with dementia tend to be isolated, excluded and subject to abuse and violence (cited in Cahill, 2018, p. 3).

The WHO Call for Action and Global Action Plan, which was adopted in May 2017, called on countries to: ‘promote mechanisms to monitor the protection of the human rights, wishes and preferences of people with dementia and the implementation of relevant legislation, in line with the objectives of the UN Convention on the Rights of Persons with Disabilities (CRPD) and other international and regional human rights instruments’ (WHO, 2016, para 20). These aims align with the principles of social work, which is committed to advocating and upholding the human rights of clients and communities (International Federation of Social Workers, 2014).

Supported decision-making is viewed as a key mechanism for delivering the rights of persons with disabilities under the CRPD. This model is founded in Article 12.3 of the CRPD and is predicated on the principle that, ‘all people are autonomous beings who develop and maintain capacity as they engage in the process of their own decision-making even if at some level support is needed’ (Devi et al., 2011, p. 254). The support model is in contrast to substituted decision-making regimes, which are systems where, ‘(i) legal capacity [the formal ability to hold and to exercise rights and duties] is removed from a person, even if this is in respect to a single decision; (ii) a substitute decision-maker can be appointed by someone other than the person concerned, and this can be done against his or her will and (iii) any decision made by a substitute
decision-maker is based on what is believed to be in the objective best interests of the person concerned, as opposed to being based on their will and preferences’ (Committee on the Rights of Persons with Disabilities, 2014, para 27). Supported decision-making acts in contrast to substituted decision-making through providing a ‘conceptual and practical bridge’ (Gooding, 2013, p. 432), which seeks to respect the individual’s will and preference, whilst viewing decision-making as an interdependent process (Sinclair et al., 2019). It allows for consideration of a disabled person’s decision-making ability, the environmental demands for decision-making and the support that is required to enable the person to decide (Shogren and Wehmeyer, 2015). The approach is informed by the social model of disability, which highlights how barriers (physical, attitudinal and structural) perpetrate disadvantage for disabled people; and feminist critiques of individualism, which explore how autonomy develops within the context of social relationships (Donnelly, 2019).

Attention has been paid to the ways in which the CRPD should be applied in situations where people are living with dementia (Keeling, 2016; Sinclair et al., 2019). However, debates remain as to how supported decision-making should be interpreted and applied in practice. Research indicates that people living with dementia are often positive about supported decision-making (Sinclair et al., 2019) although there are complex practice issues to be dealt with, particularly when the person supporting an individual may be a source of risk. Social workers are often involved in such situations and yet little analysis has been carried out on this subject, an issue that this article seeks to address.

The aims and scope of safeguarding adults’ services

Concerns about adult abuse and neglect have led to the development of adult protection systems, most notably in the UK, the USA, Canada and Australia; initially developed as a response to concerns about elder abuse in the 1980s and 1990s. A key policy document, the Toronto Declaration on the Global Protection of Elder Abuse highlighted the need for a universal human rights framework for older adults (WHO, 2002). It asserted that legal frameworks to address elder abuse were often missing, meaning that abuse might be recognised, but not adequately dealt with. Such arguments influenced responses by governments enabling the traditional focus on elder abuse to be broadened, to concepts of ‘vulnerable adults’ or ‘adults at risk’ more generally (Donnelly et al., 2017).

To some degree UK policy and law had begun this process earlier through reference to the Human Rights Act 1998. For example, the No Secrets guidance on adult abuse in England referred to abuse as, ‘a
violation of an individual’s human and civil rights by any other person or persons’ (DH, 2000, para 2.5). There remain, however, contested ideas on definitions. The term ‘adult safeguarding’ has not been defined internationally and there are differences in definitional thresholds (Mackay, 2018). Thus, all four countries in the UK explicitly state that risk of (as well as actual) harm, abuse or neglect are grounds for making an enquiry. The terminology thereafter varies: the term abuse or neglect is used in Wales and England; in Northern Ireland it is abuse, exploitation or neglect; and Scotland has the most expansive term of harm on its own. Whilst safeguarding law and policies vary across national systems, social workers tend to play a lead role in England, Scotland, Northern Ireland, Nova Scotia and British Colombia (Donnelly et al., 2017). The rationale is that social workers possess particular skills in assessment, working across professional boundaries and in enabling individuals through self-directed support. These systems have identified that they value social work knowledge. For example, The Care and Support Statutory Guidance in England states that social workers are likely to be the most appropriate professionals to make enquiries about abuse or neglect within families or informal relationships (Department of Health and Social Care, 2020, para 14.8) and highlights the importance of the principal social work role (Whittington, 2016). Nonetheless, little has been done to consider how social workers might explicitly protect the human rights of people living with dementia within safeguarding practice.

The CRPD and supported decision-making

The CRPD (Article 1) states that, ‘Persons with disabilities include those who have long-term physical, mental, intellectual or sensory impairments’, which may hinder their participation in society. This definition clearly places people living with dementia within its remit, making them subject to its rights and protections. The CRPD marks a paradigm shift for the rights of persons with disabilities as it adopts a social model of disability (identifying the need for society to adapt to the needs of the disabled person), in contrast to a medical model (focussing on cure) or a social welfare model (focussing on a person’s limitations) (Bartlett, 2012). The CRPD states that people with disabilities should be free from exploitation, violence and abuse and that state parties should take, ‘all appropriate legislative, administrative, social, educational and other measures to protect persons with disabilities, both within and outside the home’ (Article 16.1). Furthermore, those with disabilities are given positive rights and entitlements (such as the right to provision of services) by the CRPD, in contrast to the European Convention of Human
Rights, which protects individuals’ negative rights (e.g. the right to be free from undue interference or abuse from others).

National safeguarding legislation has increasingly identified the need to involve those experiencing abuse or neglect in the process (Donnelly et al., 2017), making the issue of decision-making of central importance. For example, the Care Act (2014) put safeguarding in England on a statutory footing. It is therefore essential to consider how autonomy and decision-making are conceptualised within the CRPD and how this should inform decision-making within national safeguarding practice. Protecting a person’s legal capacity and promoting their involvement in decision-making are central to the CRPD. Legal capacity can be understood as a person’s ability to hold rights, and to exercise them on an equal basis with others (Bach and Kerzner, 2010). It differs from the concept of mental capacity, which is concerned with the decision-making skills and competencies of a person, which may differ between individuals. So, from a safeguarding perspective, people living with dementia should have rights to be engaged and participate in decision-making in the safeguarding process and should also receive support to exercise these rights. Article 12 of the CRPD states that people with disabilities should be afforded legal capacity on an equal footing to others and that States should take measures ‘to provide access by persons with disabilities to the support they may require’ (Article 12.3). In English law, there is also recognition that people may have mental capacity but remain vulnerable to abuse due to manipulation or undue influence from others. In these cases, the court may exercise its ‘inherent jurisdiction’ to intervene in a way that is compliant with the CRPD (Series, 2015) (although it is beyond the scope of our article to consider the complexities of inherent jurisdiction here). Nonetheless, the CRPD Committee’s Interpretation of Article 12 identifies that people with disabilities cannot be viewed as having exercised their legal capacity unless they have been supported to decide for themselves. This view is reflected in the statement that:

State Parties’ obligation to replace substitute decision-making regimes by supported decision-making requires both the abolition of substitute decision-making regimes and the development of supported decision-making alternatives. (United Nations Committee on the Rights of Persons with Disabilities, 2014, para 28)

However, the UN Committee’s interpretation of Article 12 has been viewed as problematic by some as it states that substituted decision-making mechanisms are outlawed by the CRPD. The removal of substituted decision-making in all circumstances may cause a range of practical problems in adult safeguarding where an individual is unable to decide for themselves (as may be the case when an individual is living with advanced dementia and experiencing abuse or neglect) (Freeman et al., 2015; Gooding, 2015). No state who is a signatory to the CRPD has
followed this binary approach to decision-making in the field of capacity laws, partly because of possible, perverse outcomes that might follow. For example, the Mental Capacity Act (2005) (MCA) in England and Wales defines mental capacity in relation to decision-making and states that individuals should be assumed to have mental capacity, unless it can be established otherwise on the balance of probabilities. The MCA states that consideration of capacity must be decision and time specific. In the context of an adult safeguarding case, this means that once it has been established by the decision-maker that the person lacks mental capacity, section 4 of the MCA allows for a form of substituted decision-making by allowing the decision-maker to act in the person’s ‘best interests’. However, this places the MCA in tension with the CRPD due to its focus on decision-making capacity rather than legal capacity (Martin et al., 2016).

Current State responses to the CRPD tend to involve a hybrid mix of safeguards and processes that professionals are expected to adhere to in order to support the exercise of a person’s legal capacity (Davidson et al., 2016). In doing so, in a more limited way than the CRPD strives for, improved approaches to supported decision-making can go some way to protect the legal rights of persons living with dementia. Several arguments are presented for such approaches. First, people living with dementia will have formed a range of moral, political, social and other views before developing the condition (Donnelly, 2019). The use of mental capacity laws allows these former wishes and values to be used in preference to their current views (which may have altered radically since the onset of dementia). For example, in Briggs v Briggs [2016] Charles J. gave primacy to previously expressed wishes, in line with the ‘enabling’ ethos of the MCA deciding that, ‘an earlier self can bind a future and different self’ (para 53). As noted by Ruck Keene et al. (2017, p. 135), this can be promoted when the previously expressed and current wishes are consistent, either because they match (see Westminster City Council v Sykes [2014]) or because the person who lacks capacity is no longer able to express their wishes (see PS v LP [2013]). However, it becomes more problematic when there is a clash between a person’s past and present wishes. Domestic case law is inconsistent and the CRPD is silent on the primacy point. Ruck Keene et al. (2017, p. 138) suggest that the CRPD Committee’s interpretation of Article 12, ‘drives inexorably towards prioritisation at all points of a person’s immediately identifiable wishes and feelings’. But this approach could be problematic in a safeguarding context for persons with dementia who might express a current preference, which puts them at risk.

Second, older adults experience higher levels of abuse and neglect than other disabled groups and tend to afford greater weight to professional review and protection (Bach and Kerzner, 2010; Donnelly, 2019). This indicates the need for legal frameworks which balance notions of
empowerment and safeguarding. Such circumstances have led some to argue that supported decision-making should be the preferred option to accommodate a person’s rights under the CRPD, but that mental capacity laws are required where individuals with conditions, such as dementia may place themselves at serious risk and where there is danger in delay (Freeman et al., 2015).

Social workers are required to work within existing legal frameworks, despite the earlier stated tensions that exist between the interpretation of legal capacity identified by the CRPD and domestic laws. It is crucial that they find ways to maximise the rights of individuals to exercise their legal capacity whilst ensuring compliance with these domestic laws.

Supporting people living with dementia to take part in safeguarding decisions in England

The following section explores how supported decision-making can be facilitated in England, one of four jurisdictions in the UK. The population of England was 55.6 million in 2018 (Office for National Statistics, 2018). The most recent estimate of people with dementia, in 2013, found that 685,812 people were living with dementia (Prince et al., 2014). As in some other jurisdictions, social workers play a lead role in safeguarding and substitute decision-making processes, using a range of laws and policies, now described and discussed.

The legal and policy context for safeguarding in England

In England, the Care Act, 2014 (CA) is the key legislation for safeguarding. The Care and Support Statutory Guidance describes this as a process of, ‘Protecting an adult’s right to live in safety, free from abuse and neglect’ (para 14.7). In cases where a safeguarding referral for a person living with dementia is made, practitioners must consider their duties under section 42(1) of the CA which requires the Local Authority to consider whether there is reasonable cause to suspect that an adult:

1. Has care and support needs;
2. is experiencing, or is at risk of abuse and neglect; and
3. as a result of their needs is unable to protect themselves from the abuse or neglect or risk of it.

This process may not be linear and actions to safeguard a person may take place as part of the section 42(1) process or during a general assessment of need (LGA/ADASS, 2019a). Safeguarding decisions must be focussed on the principles inherent within the CA, notably the duty to promote well-being under section 1, and should adopt a flexible approach focussing on what matters to the individual. Decisions must also
be grounded in the six safeguarding principles contained in the Care and Support Statutory Guidance (empowerment, prevention, proportionality, protection, partnership and accountability). Workers need also to consider how abuse can be prevented (Care and Support Statutory Guidance, para 2.1) and should draw on the Making Safeguarding Personal approach. This is a sector-led initiative supported by the Local Government Association, the Association of Directors of Adult Social Services and other bodies. It promotes a personalised approach, where adults at the centre of the safeguarding process are asked what their preferred outcomes are. A number of studies suggest that these initiatives can promote increased confidence amongst staff when involving service users in decision-making (Cooper, 2015; Butler and Manthorpe, 2016). The principles of the MCA must also inform any safeguarding interventions (see further below).

**Tensions between English law and the CRPD**

The CRPD is an international treaty and therefore does not have the same status and enforceability as domestic law. Although it is not directly legally binding on the UK, it is nevertheless of persuasive authority. The Court of Appeal has affirmed the influence of the CRPD (Burnip v Birmingham City Council and Another [2012]), and there is evidence that CRPD principles are informing the jurisprudence of the higher courts, for example, in relation to decisions about the management of a person’s property and deprivations of liberty (LB Haringey v CM [2014]; P (by his litigation friend the Official Solicitor) v Cheshire West and Chester Council & Anor [2014]). However, judges have also urged caution when considering how the CRPD should shape domestic law. For example, Hayden J. noted that, whilst courts should seek to interpret and apply national laws in line with international obligations, ‘the court cannot by a process of statutory construction simply ignore or rewrite the clear provisions of the MCA [Mental Capacity Act]’ (Lawson, Mottram and Hopton, RE (appointment of personal welfare deputies [2019])). This makes it clear that practitioners must follow domestic law and cannot use the CRPD to circumvent it.

Despite the disparities between Article 12 and the substitute decision-making regime of the MCA, the MCA nevertheless has an empowering ethos and includes several mechanisms which are designed to promote autonomy and support the decision-making ability of individuals. Foremost, section 1(2) of the MCA states that individuals are assumed to have mental capacity, unless it can be established otherwise, and that they should be supported, as far as possible, to make their own decisions. The MCA (section 1, statutory principle 2) and Code of Practice (Department of Constitutional Affairs, 2007, Chapter 3) make clear that,
before deciding that someone lacks capacity, practitioners should take practical steps to help individuals to decide for themselves, including providing relevant information; communicating in an appropriate way and putting the person at ease.

The best interests checklist in section 4 includes a list of factors for the substitute decision-maker to consider. The list expressly includes the person’s wishes and feelings. Whilst they are not determinative, the court has made it clear that they must be central to the decision-making process. For example, in Wye Valley NHS Trust v Mr B [2015] the Court of Protection stated that it may in some circumstances support a person’s incompetent wishes and feelings. There is a growing evidence in the case law of the court’s willingness to engage with ‘the person and their identity’. As Series (2014) has argued, by prioritising this subjective approach to discerning best interests, the MCA can be applied in ways, which accord with the CRPD’s approach. Sections 24–26 of the MCA make provision for advance decision-making, which allows a person with mental capacity to refuse specific treatments in the future, should they lose capacity. This is regarded by the Court of Protection as a key mechanism for promoting a person’s ‘capacity to shape and control’ decisions affecting their life (Barnsley Hospitals NHS Foundation Trust v MSP [2020]). Sections 9–14 of the MCA provide further mechanisms through which people can take out a Lasting Power of Attorney (LPA). It is a legal document stating who an individual would like to manage their property and finances or health and welfare should they lose mental capacity to make such decisions. Whilst LPAs can be viewed as problematic (because they allow for decision-making on behalf of the person), they can be made to work in a CRPD context as long as the LPA holder focuses on the subjective views/wishes, etc. of the individual (rather than objective criteria) in making decisions (Series, 2014).

Supported decision-making with people living with dementia in practice

Safeguarding decisions may focus on a range of complex areas, including domestic abuse, physical abuse, sexual abuse, financial abuse, discrimination or neglect. Dementia may affect a person’s ability to make decisions about abuse or neglect and this generally becomes more severe over time (Fetherstonhaugh et al., 2013). Nonetheless, a person’s ability to decide can be enhanced through support by professionals and carers; particularly where the dementia is mild or moderate. At a practical level, supported decision-making focuses on the environmental demands for decision-making (such as consideration for the procedure in question, the physical space that the person is in and the relationship between the individual and the decision-maker). It also focuses on the support that is
required to enable the person to decide (Shogren and Wehmeyer, 2015). Currently, there are no empirically tested decision tools that have been designed to help people living with dementia to engage in safeguarding (Wied et al., 2019). However, practitioners can design strategies, based on the principles of supported decision-making, which tailor information to the needs of people with dementia and seek to involve them as much as possible in the decision-making process. In the following section, we consider how such strategies may be used, drawing on the research evidence.

In order for safeguarding to be effective, people living with dementia need to be clear what safeguarding means. This is important as the first principle of safeguarding is empowerment (Department of Health and Social Care, 2020); meaning that adults should be supported and encouraged to make their own decisions with informed consent. Such consent can only be achieved if the person with dementia is clear about the enquiries which may be made and what their options are. The MCA Code of Practice places emphasis on providing information to the person, stating that it should be tailored to their needs and ‘in the easiest and most appropriate form’ (DCA, 2007, para 3.8). This is crucial because the literature suggests that giving people with mental impairments excessive information is often challenging because of problems of cognitive retention (Wied et al., 2019). Local Authorities must therefore consider the most effective strategies for informing the public about safeguarding. Whilst the sections of the CA associated with safeguarding (sections 42–47) have been in force since 2015, levels of public awareness about safeguarding remain unclear. To ensure that people with dementia have adequate and appropriate information to make a decision, Local Authorities need to provide accessible and clear information, setting out the types of abuse, which may be experienced and how people can report it. This can be achieved by using clear and simple language with a focus on consistency of expression, as well as pictures or drawings (Wied et al., 2019). When explaining the safeguarding process at an individual level, practitioners may draw on public information as communication aids, but need to explain to individuals how it applies to them. Research indicates that people living with dementia are better able to engage where workers adopt a spirit of collaboration, highlight what they are expecting of them and work with them to define what it is they need to decide on (Groen-van de Ven et al., 2017). Practitioners should therefore explain the nature of the safeguarding concerns from the outset, identifying first how it has been raised and then what information is required.

Within any supported decision-making process, consideration should also be given to the location of the meeting. The MCA Code of Practice advises that practitioners should choose a quiet place where discussions cannot be easily interrupted (DCA, 2007, para 3.13). Current research
on this issue is limited but indicates that people living with dementia find it harder to make decisions in noisy or cluttered environments (Wied et al., 2019). Interviewing a person in a quiet room rather than a busy area is likely to improve communication. Efforts should also be made to limit the number of people taking part in an interview, particularly if they are unfamiliar to the person (Fetherstonhaugh et al., 2016).

Supported decision-making relies on building a relationship with the person. This is something that is currently overlooked in the MCA Code of Practice, which focuses more broadly on providing information and putting the person at ease (DCA, 2007, paras 3.10–3.15). In order to build an effective relationship, several factors should be borne in mind. A recent study found that persons with dementia prefer to be supported by people that they know well (Sinclair et al., 2019). Where family members are not suspected of abuse or neglect, then social workers and other professionals should engage with them so that they can provide advice on the person’s preferences and how best to involve them in decisions. Whilst people living with dementia may be fully autonomous, they may also engage in shared decision-making with carers or may delegate decision-making (Smebye et al., 2012). When people with dementia consent to these arrangements, they should be considered as ways of facilitating decision-making. In cases where it is not possible to work with family members or carers and communication is challenging, advocacy under section 68 of the CA (2014) should be considered, although this can only be provided if the conditions of the CA are met. Representation by an Independent Mental Capacity Advocate (IMCA) can also be considered where safeguarding issues arise, even where the person has friends or family (s. 4, Mental Capacity Act, 2005. It should, however, be noted that advocacy provision across England is patchy, with most Local Authorities failing to meet the spending recommendations prescribed by the Local Government Association, making person-centred practices a challenge (Dixon et al., 2020). When building relationships with the person, practitioners also need to assess their attitude to risk. Recent safeguarding guidance has placed an emphasis on positive risk-taking, in which individuals are enabled through a careful consideration of the risks in question (LGA/ADASS, 2019b). Nonetheless, research has found that people living with dementia and family carers often conceptualise risk in negative terms because of its emotive connotations (Stevenson et al., 2019). A way of dealing with this dilemma is to encourage people living with dementia to view risk in terms of ‘likelihood’ to enable positive risk-taking. Social workers should also be aware that people living with dementia may be concerned about the risks which social care services may pose to them. For example, lesbian women with dementia have been known to conceal their sexual identities because they fear discrimination by services (Westwood, 2016). Social workers therefore need to consider the person’s personal and cultural needs.
With lesbian and gay service users this may be achieved through taking account of the person’s sexual identity, making sure that it is explicitly acknowledged in safeguarding plans and through facilitating access to support networks where required.

Time is an important issue if people living with dementia are meaningfully to be engaged in decision-making. The MCA Code of Practice places emphasis on the timing of conversations, stating that decisions should not be rushed and that unnecessary time limits should be challenged where the decision is not urgent (DCA, 2007, para 3.14). This guidance is supported by research which has found that supported decision-making processes are more likely to be effective where a person living with dementia is given time to recognise the issues they face and consider the options to enable a final decision to be made (Smebye et al., 2012; Fetherstonhaugh et al., 2013). Ideally, time should be ring-fenced, to enable an assessment of the person’s life story, and conducted at a pace that they feel comfortable with and at the time of day during which they function best. These recommendations are congruent with guidance by the Local Government sector (LGA/ADASS, 2019a,b), which has encouraged practitioners to view safeguarding as a series of conversations with the person, drawing on a strengths-based approach. There are possible organisational impediments to these aspirations where resources are limited. In some instances, however, local authorities have supported a flexible approach. For example, the London Safeguarding Adults Board (2019) states that a divergence from target timescales may be justified for a number of reasons including the need to provide supported decision-making. Nonetheless, there may be situations where immediate risks prevent engagement with the person over time, discussed in more detail, below.

Practitioners should design strategies that tailor information to the needs of people with dementia. As mentioned above, there are no empirically tested decision-tools to enable clients to engage in safeguarding (Wied et al., 2019). The MCA Code of Practice, however, provides guidance on what steps can be taken to tailor the information to the individual and ensure it is ‘relevant’, including not giving too much detail; providing a ‘broad simple explanation’ and outlining the risks, benefits and effects of the decision (2007; para 3.9). It has been found that strategies which build a relationship with the person through helping them to feel useful and productive are most effective (Fetherstonhaugh et al., 2013). At a practical level, this involves writing options down, to ensure the retention of information; the use of lists to explore options and using visual aids (such as pictures or photographs) to compensate for memory problems. Limiting decisions to two or three options to prevent the person experiencing ‘sensory overload’ has also been found to be important (Smebye et al., 2012; Fetherstonhaugh et al., 2013). However, this option needs to be considered carefully. Not giving the full range of options
may lead to over-simplifying or withholding important information. This is problematic from a legal perspective as it limits how informed the decision can be, thereby impacting on the person’s rights. When deciding how to proceed, workers need to consider the person’s individual preferences for decision-making as well as the potential consequences of the decision. Further sources of support from family/friends or professional advocacy services should be considered as a way of maintaining the person’s legal capacity, as recommended by the MCA Code of Practice (DCA, 2007, para 5.69).

Consideration needs to be given to principles of safeguarding where an urgent decision needs to be made. Whilst the MCA makes no explicit reference to safeguarding, it aims to balance an individual’s right to make decisions with ‘their right to be protected from harm if they lack capacity’ (DCA, 2007, para 1.4). Relying solely on the concept of mental capacity may not accord with the approach to legal capacity within the CRPD, but can be viewed as necessary in cases where a person with a mental health problem is at serious risk and there is danger in delaying decisions (Freeman et al., 2015). Whilst section 4 of the MCA allows for a best interests decision to be made, the person’s legal capacity can still be protected where workers are able to draw on advance decisions, designed to attend to previous choices made by the person (Series, 2014; Keeling, 2016). In order to maximise legal capacity, these should be referred to first, although in practice their use is likely to be limited, as they focus on advance refusals of medical treatment. Where neither an advance decision, a LPA, or a court-appointed deputy exists, practitioners need to resort to a best-interests decision-making process in line with the MCA, although this should be viewed as a last resort after all other decision-making avenues have been explored. To maximise the person’s rights, all efforts should be made to consider the subjective wishes of the person within this process. In these circumstances, practitioners should endeavour to resume supported decision-making once the person is out of immediate danger.

Finally, it should be noted that there are some limits to the research evidence as it stands. Although the CRPD has led to an increased emphasis on supported decision-making, research on supported decision-making remains at an early stage, particularly with regards to dementia (Wied, 2019). Whilst current research may inform practice, many of the studies focus on aspects of supported decision-making, such as user-involvement or participation, rather than on the supported decision-making process as a whole. It should also be noted that much of the existing evidence draws on qualitative research. Whilst such research has provided valuable insights, there is a need for studies that test the effectiveness of supported decision-making for people with different types of dementia. Such developments have the potential to lead to empirically tested decision-making tools with greater levels of validity.
Conclusion

Dementia leaves individuals more susceptible to abuse and neglect and action is required to address this. Social workers play a key role within adult safeguarding systems internationally and have an opportunity to address such abuse, yet little analysis has been carried out on this issue. The CRPD provides social workers with the opportunity to strengthen human rights protection for people with dementia, through the application of supported decision-making. This opportunity should be welcomed whilst recognising practice dilemmas, particularly in navigating the tensions between international frameworks and domestic law. In England, these are illustrated by the CRPD’s insistence on supported decision-making in Article 12, compared with the MCA which embraces a substitute decision-making model, albeit with elements of supported decision-making built into the process. Despite these tensions, steps can be taken to maximise the legal capacity of people living with dementia to promote the ethos of the CRPD through adopting a range of supported decision-making strategies.

Social workers must adhere to the provisions in the MCA and take appropriate steps to aid decision-making ‘before’ an assessment of mental capacity is made, in line with the guidance in the MCA Code of Practice. Additionally, key measures should also be taken to maximise supported decision-making. Local authorities should be required to provide clear and accessible information to the wider public and to people living with dementia, their family and carers. These should explain what safeguarding is, how the safeguarding process works and how to access it. It is imperative that social workers clearly explain to the person how the safeguarding concern has been raised, associated issues and what information they require. Drawing on the research evidence, it has been argued that key steps are involved in good quality safeguarding interventions that are service user focussed. For example, people living with dementia should be interviewed in quiet areas, with care being taken to minimise the number of attendant people in the room. Skills in building effective relationships are critical for the practitioner. People living with dementia prefer to be supported by people that they already know, although advocacy under the CA should be considered where this is not possible and advocacy under the MCA may be considered where the person lacks capacity. Social workers also should be mindful that people with dementia often frame notions of risk differently to that of professionals. This awareness and knowledge can help build on the preferences of the person. The importance of time is central to decision-making processes. Ideally it should be ring-fenced in order to learn about the person’s life story, and to enable assessments to be conducted at a pace that the person feels comfortable with, and at the time of day at which they function best. The use of visual
aids, diagrams and lists have been shown to assist the person to retain information and make decisions. Consideration should also be given to limiting options to enhance comprehension. Where immediate risks prevent this, the person’s capacity should be protected through ascertaining wishes expressed in advance decisions or through an LPA, or court appointed deputy, where they exist.

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