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Is personalisation the right plan at the wrong time?: Re-thinking cash-for-care in an age of austerity

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Abstract
As personalisation has dominated the policy agenda for reform in social care, its roll out as a mainstream option has coincided with global programmes of austerity which have targeted services for disabled people. Was this simply bad timing or was its implementation always part of the agenda for reform? Whilst the principle of cash-for-care schemes drew strongly on promoting the independence and autonomy of disabled people, early incarnations of direct payments policy in the UK were initially at least in part rolled out in light of potential cost savings to social care budgets. This article explores these and other issues in light of evaluations of the Scottish Government’s self-directed support test sites (2009-2011) and implementation of subsequent legislation enforcing models of self-directed support in Scotland from April 2014. Discussion draws on the authors’ recent book (Pearson, Ridley and Hunter, 2014) to reflect on what the evidence can tell us about radically transforming social care at a time of global austerity.

Keywords
Personalisation; self-directed support; austerity; co-production; Scotland; disability

Introduction
Over the past 20 years, personalisation has emerged as a reforming concept across many areas of social policy in the UK (see Fox et al, 2013 for criminal justice; Pykett,
2009 for education; Toerien et al, 2013 for employment; Forder and Jones, 2014 for health; and Pykett, 2009 for education), gaining what West (2013) identifies as an ‘ideological grip’ across the public sector. Its appearance in UK systems of social care draws in part from the legacy of disability activism culminating in a long fought campaign to secure direct payments as a facilitator of independent living (Campbell and Oliver, 1996; Morris, 1993), alongside a drive to ‘modernise’ and provide more cost efficient and flexible modes of service provision in emerging local care markets (Pearson, 2000; Spandler, 2004). Accordingly several commentators observe that personalisation has come to stand for a number of competing and contradictory demands: perceived as either a radical transformation of social care wherein the power balance shifts in favour of service users, or as an attempt to roll back the boundaries of the welfare state, to introduce neo-liberal ideals, and undermine public sector services and responsibilities (Needham and Glasby, 2014).

Elsewhere European models of personalised social care have emerged with a similar diversity in policy discourse, varying from the rights-based scheme of Sweden, to more restrictive approaches – for example in Belgium, France and Germany – where policy set out specifically to reduce the numbers of persons entering residential care (Glasby, 2013). Yet across the UK (Slasberg et al, 2015), Europe (see Hauben et al, 2012) and North America (Power, 2014), the impact of austerity on personalised support in social care has been acute. Despite its roots in the independent living movement (Oliver and Sapey, 2006), budget cuts and the broader reconfiguration of support for disabled people has acutely undermined these principles. The impact of austerity not only reflects a trend in disability activism, but one that has spread across services for other user groups. For example, Roy and Buchanan’s (2015) recent work
shows how uncertainty and diminished budgets have greatly reduced investment in user activism through recovery services. Consequently, they argue that the parameters for providing services with social value, utility and effectiveness - the corner stones of independent living - have become far more limited. In recent years, commentators such as Taylor-Gooby and Stoker (2011) have positioned the UK programme of austerity reforms as part of a broader restructuring of state services with a key goal to transfer responsibility from state to the private sector and ultimately to citizens. In these terms, others have identified personalisation as the neo-liberal tool to facilitate this process across social care (Ferguson, 2007; West, 2013), signifying the end of collective services and community based support.

Whilst Scotland has traditionally been slow to embrace the market in social care and was largely sceptical of previous policy incarnations of personalised support (see, 2004), its implementation of new legislation for self-directed support (SDS) undoubtedly takes social care into a new era. In this article, we explore the promotion of this new era for personalisation in Scotland at a time of austerity. We begin by highlighting the challenges of the personalisation agenda in a broader European context, as many models of personalised support have also been compromised by acute budgetary cuts. Discussion then moves to a commentary on Scotland’s positioning of SDS in its early stages of implementation. This provides a more in-depth focus, by drawing on a series of findings from the evaluations of SDS test sites in three local authorities across Scotland (Manthorpe et al, 2011; Ridley et al, 2012; 2014), information from the roll-out of SDS across Scotland (Scottish Government, 2013) and data from a Freedom of Information (FOI) request sent out to all 32 Scottish local authorities in April 2015. This asked authorities to indicate how many
users, across different groups, were in receipt of each of the four SDS options. We consider whether the early roll out of SDS offered a full range of personalisation choices in Scotland. In doing this, we ask whether it is in fact the right plan for reform to facilitate independent living, but whose legitimacy has been weakened amidst the current spending restraints. Alternatively, was its role in the dismantling of local networks of community social care services, loss of community spaces and reducing access to personalised support an inevitable consequence of a neo-liberal agenda for welfare reform?

**Austerity and European challenges to personalised support**

Before looking at changes in Scotland, we turn to briefly to look at how personalisation policies have fared across Europe in response to the hostile economic climate. As the European Agency for Fundamental Rights warned in 2012, the long-term impact of austerity changes clearly has serious implications for the well-being of a generation of disabled people and would progressively erode the advances that have been made in establishing and promoting the rights of disabled people (Hauben et al, 2012). In their review of evidence of how austerity measures have impacted on the rights and status of disabled people, austerity has undoubtedly emerged as a negative influence, with significant differences over its impact on individual disabled people’s lives observed across Europe. For example, countries such as Germany, Austria and Scandinavia appeared to have endorsed far fewer reductions in social benefits and social services, particularly in contrast to Greece, Ireland, Spain and Portugal. Like the UK, Hungary has tended to frame austerity measures through a broader programme of welfare reform and discourse of modernisation. Yet overall, the pattern
indicates a clear focus on the reduction of social services, thereby disproportionately impacting on disabled people (Pearson, Ridley and Hunter, 2014).

As we detail later in this article, access to a personalised budget has become increasingly restricted as the austerity cuts have taken hold. This is a trend that is also being replicated across Europe. Even in member states with well-established independent living schemes, key changes have been made which limit disabled people’s access to personal assistance. One of the tools used to restrict access has been through use of waiting lists. In Ireland, for example, waiting lists for social and health care have sharply increased and more than 25 percent of persons with physical impairments are waiting for access to assessments for PA and support services (Hauben, 2012). Similarly amongst Belgium’s Flemish community, the PA waiting list has increased to 50,000 people - resulting in an estimated 5-10 year wait for this type of support (Ratza, 2012). In other countries such as Greece, where there are no PB schemes in place, financial support to enable disabled people to live more independently has simply been cancelled. Likewise in Portugal, plans to introduce PBs from 2011 have been delayed indefinitely (Hauben, 2012).

**Austerity, personalisation and a new era for social care in Scotland**

Like many of the examples shown across Europe, for local authorities across the UK, rolling out of personalisation and SDS in social care has also coincided with the economic crisis. In Scotland, the Social Care (Self Directed Support) Act 2013 was implemented from April 2014, just as many of the austerity measures were beginning to take force. Leading up to this period, overall Scottish spending was reduced by around 11 percent in real terms across a four year period, leaving a 7 percent cut in
resources across local authority spending and 37 percent in capital spending (Scottish Government, 2012). As social care forms a core service within local authorities, cost cutting of local budgets has acute implications for its users. Whilst this had already had a significant impact on the planning and delivery of services (Main, 2013), it was clear that changes were on-going. Indeed analysis by the Scottish Government indicated that the worst of the cuts were yet to emerge – with the full impact of changes expected in 2016 (Scottish Government, 2014). Furthermore, the election of a majority Conservative Government in May 2015 with a commitment to cut welfare spending by a further £12 billion signified a longer term decline for funding in these service areas (Watt, 2015). As we show in this article, front line social care workers in Scotland have continually expressed strong concerns about the current and future impact of austerity measures in adopting SDS. In order to examine these issues in more detail, we draw on findings from evaluative studies of SDS as the basis of a critical commentary on the positioning of policy in the early stages on implementation. The following section details the background and methodology to this work.

**Background to the test sites and methodology**

In 2009, as government cuts on social welfare spending were starting to impact, the Scottish Government, Convention of Scottish Local Authorities (COSLA) and the Association of Directors of Social Work (ADSW) selected three local authorities to design and test out the systems needed to implement SDS in social care. The test sites were selected to represent a broad geographical spread including one remote rural site, and local authority performance based on Social Work Inspection Agency inspections. Each test site was given up to £1.2 million for just over two years (January 2009 to 31 March 2011) to put in place systems that would facilitate a shift towards SDS. At the
outset, the Scottish Government wanted the test sites to focus on three themes identified from research evidence as critical to enable this transformation: these were the need to reduce bureaucracy or ‘red tape’; the importance of leadership and training; and the need for bridging finance to double fund existing services while introducing more individualised models of support.

The evaluative studies that we draw upon for this article, used a mixed methods approach involving several elements. The first evaluation comprised a review of the literature on facilitators and barriers to SDS (Ridley et al 2011); gathered and collated quantitative information about those receiving SDS packages; conducted secondary analysis of national statistical data (Scottish Government, 2011); carried out interviews with national and local stakeholders; held action learning sets involving key stakeholders from each area; monitored test sites’ implementation plans; and focused on 30 individual service users’ and carers’ experiences to better understand these new SDS systems from service users’, family carers’, and front-line workers’ perspectives.

A follow-up evaluative study involved interviews with key stakeholders, a questionnaire survey of frontline staff and gathered systematic monitoring information to provide an update on developments in terms of the processes, practice, outputs and outcomes of SDS in the former test site areas (Ridley et al, 2012). The views of over sixty different stakeholders from the three areas were analysed including service user organisations, advocacy services and voluntary sector providers. Over 500 frontline staff in adult social care across the three areas were surveyed via an online questionnaire (with a response rate of 43 per cent). This article draws in
particular upon the findings from this survey to reflect upon frontline workers’ experience and perceptions of the impact of budget cuts on SDS implementation.

In order to provide an updated snapshot of SDS implementation in Scotland, we also carried out a FOI request across the 32 local authorities in April 2015. Authorities were asked the question: what are the total numbers of users currently receiving SDS for each of the four options? At the time of writing, 29 local authorities had replied. Twenty six of these offered full information on the questions asked, three only included partial data and two local authorities declined the request on grounds of cost. The implications of these findings are discussed later in this article.

Ethical and research governance approvals were received for the research from [Authors’ University ethics committee] as well as from the ADSW and the three local authorities.

In this article, we consider findings across these studies alongside a FOI request to explore how the timing of the austerity cuts has impacted on the implementation of SDS. This is discussed through four key issues. Firstly, by looking at the shift away from direct payments to SDS, we consider how grassroots disability activism and its considerable influence in the development of direct payments became diluted under the infiltration of neo-liberalism and the personalisation of social care. Secondly discussion moves to examine these ideas more specifically in the Scottish context by looking at how personalisation has been adopted through new legislation and what we define as an ‘evolving concept of SDS’. Thirdly we return to issues relating to the discourse of cost efficiency associated first, with direct payments and then with the broader personalisation agenda. This has clearly been a significant theme in the
adoption of personalisation policies across the UK – particularly when comparing to other countries such as Norway. Indeed, we show how findings from the test sites offer some evidence to support this contention, whereby in some areas, SDS has been used as a mechanism to cut support packages. This links into the final area, a focus on the limitations and problems of the resource allocation system (RAS).

From DPs to SDS

As has been documented extensively elsewhere (e.g. Campbell and Oliver, 1996; Glasby and Littlechild, 2009) the idea of cash-for-care as an alternative to directly provided services originated in the 1970s in the US (Arksey and Kemp, 2008). This saw a small group of students with physical impairments at the University of Berkeley secure payment of cash in place of services to buy-in support which fitted their daily needs. Gradually news of this spread throughout the US and by the 1980s, similar challenges were being made by groups of disabled people in the UK. Throughout the 1980s and 1990s indirect payments – those administered through third parties such as voluntary organisations – were used to overcome legal restrictions to a cash-based model of support across the UK. In line with actions a few years earlier in the US, the roles of small groups of disabled people were central to changes during this time. They demanded a more flexible alternative to rigid and paternalistic modes of service provision offered to them by local authorities. In Scotland, although developments were more limited, pockets of activism also emerged during the 1980s, notably in the former Lothian region. At this time, Lothian Regional Council permitted three indirect payments to be made. While this was initially approved, problems arose when, in 1995, the then Department of Social Security was alerted to this practice and suspended all the payments of those involved. A challenge to the case in Lothian was
made and won by the local authority, but the case had raised the profile and potential of cash payments to increase disabled people's choice and control over their social care, thereby instigating the need for a formal legislative path to be developed (Pearson, 2004). Disability activism therefore undoubtedly played an important role in moving towards legislation for direct payments in the late 1990s, although as we discuss shortly, a discourse of efficiency always prevailed.

Whilst many welcomed the shift to direct payments, change was not uniformly embraced with strong resistance emerging in many parts of the UK. In particular, the number of users in Scotland, Wales and Northern Ireland were especially low (Riddell et al, 2005) when compared with key local authorities in the south of England such as Hampshire and Essex. From 1997-2003, various attempts were made by the then Scottish Executive and Westminster Governments to reignite policy but uptake was never more than marginal (Priestley et al, 2010).

Under the Labour Government (1997-2010) many of the themes which linked direct payments to the Conservative's broader agenda of marketisation were developed through a focus on the personalisation of social care services (Pearson, Ridley and Hunter, 2014). At this stage, ideas promoted by Leadbeater (2004; 2008) exploring how services could be modernised through the direct participation of users in service delivery, had a profound influence on reforming the adult social care system. Leadbeater et al (2008: 47) argued strongly in favour of a model of consumerism which encouraged users to take on the role of a budget holder, but at the same time embracing themes of citizenship and related notions of rights and entitlements.
Although widely popular amongst Westminster government circles from the mid-2000s, caution was expressed from a number of quarters.

For many, the promotion of consumerist identities sat uncomfortably with the experiences and identities of many people who use social care and mental health services (Carr, 2011; Morris, 2004). Others such as Ferguson (2012) and Dodd (2013) argue that the social justice values associated with personalisation policies and which were so strongly promoted by the disability movement, have been subsumed by neoliberalism. Therefore rather than extend individual autonomy, Ferguson warned that such policies would be used by cash strapped local authorities to make savings from their services. As we discuss shortly, it is clear that this neo-liberal discourse has remained dominant in the roll-out of personalisation and is an unhelpful legacy for its implementation (Pearson, Ridley and Hunter, 2014).

In Scotland the shift to a more personalised system of social care has been slower and more complex (Kettle, 2011). As stated, this undoubtedly reflected a less enthusiastic drive towards the marketization of social care than south of the border and a reluctance in many areas to fully embrace direct payments (Pearson, 2004). As policy emerged, the Scottish Government employed the term Self Directed Support (SDS) to develop distinctive policy goals (Pearson, Ridley and Hunter, 2014) setting out SDS to be part of its aspirations to create a ‘healthier nation with stronger and safer communities’ (Scottish Government, 2007: 2).

It is clear from the history of direct payments, personalisation, individual budgets, SDS and other cash-for-care based schemes in social care that terms can be used
interchangeably and are confusing (Pearson, Ridley and Hunter, 2014). Notably direct
payments and personal budgets have often been set out as being essentially the same
thing – in that they both involve a cash alternative to directly provided services.
However as Beresford (2009) notes, there are important differences in their
underpinning ideologies. As discussed earlier, direct payments were a grassroots
development emerging from the disabled people’s movement and a desire to equalize
opportunities and increase independent living (see Glasby and Littlechild, 2009),
whilst personal budgets evolved from developments largely driven by professionals
critical of the welfare state and its ability to promote independent living. Direct
Payments are based on a social model of disability and the philosophy of independent
living, whereas ‘personalisation’ emerged from the vision of welfare reformers such
as In Control – pioneers of SDS and Individual Budgets in 2003 - as an approach to
enable disabled people to direct and take charge of their support and which has since
underpinned UK-wide policies promoting personalisation (Pearson, Ridley and
Hunter, 2014). The following section explores this in relation to the development of
SDS in Scotland.

**SDS - an evolving concept**

The Care and Support (Self-directed Support) (Scotland) Act 2013 (the SDS Act)
offered four different options for SDS. These included option 1, a direct payment
approach; option 2, a selection of support by the user but arranged through the local
authority or other agency on their behalf; option 3, support organised and provided by
the local authority; or option 4, a mixture of these options (Scottish Government,
2014). Critically, support for SDS from the governing Scottish National Party brought
policy into the mainstream of social care provision and thereby increased the numbers of people directing their own support from the time of implementation in April 2014. Unlike previous policy incarnations, the focus on direct payments as only one variant of the SDS approach has caused some confusion with SDS becoming an ‘evolving’ or ‘elastic’ concept, which has been variously defined and implemented much as has personalisation (Manthorpe et al, 2011; Larsen et al, 2013). As discussion moves to look at some of the broader policy discourses, we will re-visit this theme later in the article.

Commentators identify a perceptible shift in the rhetoric around personalisation, moving from a primary focus on individual control over budgets, to an emphasis on the importance of offering ‘choice and control’ and focusing on individual outcomes (Beresford, 2009). The experience of the test sites indicates a similar evolution of the concept of SDS in Scotland. Early on, the stated purpose of the test sites had been to increase the number and range of people receiving direct payments:

“The very baseline is Manifesto commitments, each of the test sites must demonstrate increase in take-up of direct payments as an absolute …” (Interview with national policy makers in 2010 quoted in Pearson, Ridley and Hunter, 2014: 31-32).

In contrast, local and national stakeholders insisted that SDS was a broader concept implementing ‘personalised services’, and that it consisted of a spectrum of options that included direct payments at one end and local authority-provided services at the other. This broader definition of SDS was also in evidence in the National Strategy
for SDS (Scottish Government, 2010), as well as in the subsequent SDS Act. Furthermore, the range of support packages evidenced during, and a year after the test sites, reflected this. Initially at least, direct payments (option 1) was the most common SDS option (Manthorpe et al., 2011), while at the follow up evaluation, the majority of SDS packages were arranged and managed by local authorities (option 3) (Ridley et al., 2012): a pattern mirroring the experience of Personal Budgets in England, the majority of which were taken in the form of local authority managed budgets (ADASS, 2011; Beresford, 2013; Slasberg et al., 2013). Indeed, by the time of our FOI request in April 2015, 73 percent of the local authorities who responded (n=26) indicated that option 3 - support organised and provided by the local authority - was the most commonly used form of SDS. In only one local authority did direct payment recipients (option 1) outnumber those in option 3.

**A discourse of cost efficiency**

Since legislation implementing direct payments in the late 1990s, and the promotion of personalisation by Leadbeater et al (2008), a discourse of cost efficiency has been a major feature of policy promotion across the UK. Even before this time, the successive dismissals of direct payment legislation on the statute throughout the 1990s were only brought to an end when the British Council of Disabled People commissioned research, which in promoting the merits of direct payments also showed them to be up to 40 percent cheaper than directly provided services (Zarb and Nadash, 1994). Whilst this has never been realized in practice, the link between personalisation and cost efficiency has remained a dominant theme in the UK. Leadbeater et al (2008) and others (see Duffy, 2010; Needham, 2011a; 2011b) have highlighted the link, suggesting that savings could even be as high as 45 percent. This
Despite the national Individual Budget (IBSEN) study (Glendinning et al, 2008) finding an 11 percent increase in infrastructure costs thus contradicting the claim that the process could increase value for money.

It is therefore clear that the focus on budgets and empowerment through cash-for-care has been a key feature of personalisation in the UK. This is reiterated by Christensen and Pilling (2014) who examined the differences between personalisation policies implemented in social democratic welfare regimes and those in more neo-liberal contexts. In looking at this issue, they provide a comparative analysis of Norway and England and argued that whilst both ran systems that had been supported by active disability movements and promoted market based models for social care provision, the main difference lies in their promotion of care markets and the focus on consumer roles. In England a key policy emphasis has been to give users knowledge of the amount they have to spend (indicative budget) on their support upfront and to have choice and control on how this is spent (Glasby and Duffy, 2007).

Thus the speed at which IBs have been implemented in the UK was, as Carr (2013) has observed, made on the basis of scant evidence. In Scotland, the results of a small-scale IB demonstration project in North Lanarkshire were positive about the benefits and the potential of the In Control approach (Etherington et al., 2009), and this was followed by test sites adopting a similar approach in 2009 (Manthorpe et al 2011). The nexus of control promoted at the heart of the personalisation strategy in England has often been presented as entirely positive and that if people have choice and control simply by virtue of knowing the allocated budget, its actual size is unimportant. As Slasberg et al (2012, 2013, 2015) observe however, this fails to
acknowledge that positive results were associated with PBs being taken as direct payments, and further that those receiving local authority managed services did not experience the same positive outcomes. Furthermore, better individual outcomes have been found for those using a direct payment to employ PAs and have enough resource to meet social and leisure needs (Slasberg et al, 2015). By contrast, Norway’s focus has been on the control (or what Christensen and Pilling (2014) term ‘citizen choices’) they have over the actual support received, that is, the role of managing their support workers.

The experience of the test sites offers a different insight regarding upfront allocations. While not disputing that a resource allocation in the form of an IB should be transparent, the Dumfries & Galloway test site concluded that such transparency sometimes resulted in inferior solutions as it shifted the focus to the budget entitlement instead of on the outcomes sought. Instead, they concluded that prior to introducing a budget figure, identifying and working with natural support networks led to the development of more creative solutions based on individuals’ assets and community resources:

“The In Control 7-steps didn’t work for us. It was the money upfront bit. What we learnt from the test site was that if you tell people what the money is you get a plan that is that to the penny. We did a few where we didn’t say what the indicative budget was and quite a few came in at less and they were more outcomes focused. Giving the figure upfront we found was skewing people’s thinking, and the plans were not quite as creative”
As other research has shown (Hatton and Waters, 2013), user and carer satisfaction with SDS has been linked with agreement about the panel’s decision on the size of the IB in relation to what they identified was needed (Manthorpe et al, 2011). User and carer dissatisfaction in the test sites was commonly associated with there being some discrepancy between what they perceived as their needs and the final financial allocation decided by the local authority, as well as with differences between users, carers and professionals in their perceptions of need. Some users felt they had been encouraged to have ‘false expectations’ as one disabled man in his 70s demonstrates:

“They (social workers) took everything into consideration that we were telling them and they wrote it down and then they checked it against what we had written down, the form we had filled in, and they seemed quite in agreement with what we needed and what we had self-assessed and then as I say it went away and sat in front of this Board (panel) for a year and when we got word back, everything had been cut, we only needed so many hours and ... in the morning and so many hours at night for getting ready for bed and whatever...” (Quoted in Manthorpe et al, 2011: 57)

Frontline workers from the Glasgow test site in particular linked SDS with making cuts in social care budgets, identifying negative impacts on service users’ choice and control. Personalisation they said, was a ‘means to cut services’, a ‘guise to reduce
budgets’, and the process was overtly ‘budget orientated’ rather than driven by an ethos of independent living. The direct impact of cuts on social care and the inability then to implement SDS in the way intended, was a common theme, even though many typically prefaced their comments with ‘personalisation is a good idea but...’. Some highlighted increased stress for service users and carers with many experiencing a reduction in traditional packages of care when they were moved onto SDS, which impacted negatively on their quality of life. The majority of frontline respondents made this clear link:

“The SDS process is being used as a mechanism for managing budgets. However this is in relation to the overall financial climate and the cuts would have been required regardless of the implementation of SDS...Support packages tend to only deal with basic need and no longer take into consideration quality of life and in some cases the panel does not account for carers’ stress within the support package”. (Frontline worker responding to the online survey, 2012)

It is likely that such dissatisfaction with SDS budgets will continue to increase in the future, given the financial constraints that local authorities will increasingly face (see Scottish Government, 2014).

**Limitations of Resource Allocation Systems (RAS)**

One of the most controversial elements of the personalisation programme in social care has been the system for allocating resources or Resource Allocation System (RAS) (Slasberg et al, 2012; Series, 2014). The RAS is defined as ‘a set of rules that
can be used to calculate an individual budget’ (Duffy, 2010: 206). It has therefore been central in how local authorities make decisions about individual packages. Yet growing evidence suggests that RAS are operating in ways that were not anticipated by those who promoted them (Series, 2014). Indicating the strength of feeling evoked by RAS, a user led organization consulted during the test sites evaluation observed “we [disabled people’s movement] fought for SDS and personalisation but not for the RAS” (Pearson, Ridley and Hunter, 2014, 36). All three test sites struggled with setting up RAS under the In Control model, and had not resolved these issues one year on. A major limitation to RAS was found in applying across client groups:

“Getting the resource allocation right is an art not a system — it needs to be flexible. Slotting people into boxes is not going to work. At the end of the test site and now there is still a need to develop an equitable way of distributing and managing resources. We know what we have is not quite right but we haven’t solved this issue yet”. (Local authority implementation staff, Pearson, Ridley and Hunter, 2014: 59)

Dissatisfaction with RAS has tended to centre around increased bureaucracy and questions have been raised as to what constitutes a ‘fair allocation’ (Slasberg et al, 2013). As West (2013) found in her study of personalisation in an English authority, amid austerity cuts of around 30 percent, resource restrictions meant that for many, the very possibility of obtaining a PB was denied. Yet rather than address these issues directly, the Council moved forward with the scheme and presented it as part of a ‘transformation strategy’ which would help widen the social care market and drive out perceived inefficiencies in service provision. West’s work highlighted how
populations of social care users have been left outside new systems of personalised support by reframing the boundaries of eligibility. In Scotland similar patterns are beginning to emerge. For example in Glasgow, the initial development of SDS coincided with an 11 percent cut from the £89 million social care budget during 2012/13 (Main, 2013). In explaining the cuts, the local authority have presented personalisation as a strategy for services to be delivered more efficiently. It will be interesting whether defining eligibility determination using national eligibility criteria as under the Care Act 2014 in England, will alter this picture south of the border (Care and Support (Eligibility) Regulations 2014).

Research finds a major discrepancy overall between indicative and final budget figures agreed by RAS panels (Series and Clements, 2013: Slasberg et al., 2013). This was borne out to some degree by the experience of the Scottish test sites, though our data is derived from small participant numbers and is not strictly comparative with these other studies. Nonetheless, our data indicates the emergence of similar concerns in Scotland, with only a minority of frontline workers suggesting that outcomes from SDS were more positive:

"We are clearly advised to work towards less rather than more and we know before we discuss cases that the budget needs to come in less than the estimated budget and definitely less than existing service provision unless there is compelling reasons to argue differently…" (Frontline worker responding to online survey, 2012).
One year on from the test sites, frontline workers from Glasgow commented that final budget levels were invariably lower than those proposed, and that service users were experiencing cuts in support. Around three out of five agreed that SDS budgets were insufficient to meet needs, which was similar to the findings of the UK-wide Community Care and Unison PB survey (2011), showing 48 percent of social work practitioners assessing PBs as insufficient to meet service users' needs. Whilst the strength of feeling linking SDS with cuts was less pronounced in two of the test sites, frontline workers in Highland expressed disquiet that budget constraints made it difficult to get funding approval for more ‘aspirational’ rather than ‘critical’ needs, indicating a shift in the operation of eligibility criteria. Such findings strike as ironic given personalisation was initially promoted as ‘aspirational’ (Needham, 2011).

**Discussion and conclusions**

As Scotland has moved to implement SDS as the mainstream framework for social care, this article has raised a number of concerns relating to the timing and underpinning discourses influencing this shift. Findings from SDS test sites in Scotland and more recent figures on SDS take up from local authorities indicate that SDS has moved away from direct payments to a focus on support which is organised and managed by the local authority. At best, those who have entered the SDS system in Scotland may have experienced greater involvement in the discussion about their needs and support packages. However, as test site results show, there is also evidence that reassessment through a new SDS process falls short of co-production and may at worst, amount to reduced support packages. Those implementing SDS continue to face a number of thorny issues not least with systems of allocating resources which tend to become heavily bureaucratized and act as mechanisms for delivering cuts
(Beresford, 2013). As Stack (2013) observes, the reality of RAS is ‘more opaque’ than intended. The root of the problem may however lie more fundamentally as Slasberg et al (2013) argue in what is meant by ‘fair allocation’.

Furthermore, as Hall (2011) argues, the new ‘care market place’ for personalisation has both transformed existing relations of care and constrained collective and interdependent forms of care that many people value resulting in a ‘lack of fit’ between the needs of different groups of disabled people and personalised care provision. As local authorities devolve their budgets to the individual, collective forms of support will inevitably receive less resourcing (Needham, 2013). The implications of more privatized experiences of social care support and reduced opportunity for social interaction linking with broader critiques of neoliberalism across the social sciences (Dodd, 2013; Verhaeghe, 2014). As Monbiot (2014) suggests, ‘the market was meant to emancipate us, offering autonomy and freedom. Instead it has delivered atomization and loneliness’.

Whilst the cuts from local authority budgets have undoubtedly impacted on the roll-out of SDS, the establishment of the Scottish Independent Living Fund (SILF) in July 2015 – with a £5.5 million investment from the Scottish Government to continue with existing payments - represents an important departure from policy in England and Wales (Northern Ireland is also covered by the SILF). Since its inception in the mid-1990s, the Independent Living Fund (ILF) allowed an alternative route to cash payments for disabled people with the highest support needs, as funds were paid via a charity and funded through central government rather than local authorities. It proved highly popular with its users and played an important role in the evolution of
personalised support (Morris, 2004). However the UK Coalition Government (2010-2015) set in place a decision to close the ILF in July 2015 and transfer monies to local authorities in England and Wales without a requirement for them to be ring-fenced. A move that has been widely criticized and raised questions about how already cash strapped authorities would cover the fund’s costs and enable its users to sustain the same levels of independence (White, 2013). The contrast in responses to dealing with the future of the ILF north and south of the border are therefore important when looking at the broader personalisation agenda as each have very different implications for the goals of autonomy, citizenship and control.

Despite growing critiques of personalisation, Beresford (2014) argues that innovative grassroots developments have much to offer. Notably, initiatives such as those led by The Standards We Expect consortium (Joseph Rowntree Foundation, 2011) have set out a vision for person-centred support and developing ‘bottom-up’ ways of challenging barriers to this. Although highlighting the difficulties in achieving personalised outcomes in a hostile economic climate, these commentators also argue that the funding shortfall is intrinsically linked to the continued existence of a social care culture that remains at odds with person-centred support and the principle of self-determination. The solution, argue Slasberg and Beresford (2015), lies in learning the ‘right lessons’ from the implementation of personalisation so far, moving from consumerist notions of individual control and a preoccupation with PBs, to refocusing on truly personalised support plans built from accurate assessment of needs, sufficiency of resource and flexibility of provision.
In Scotland, examples of positive practice in SDS are emerging. Research funded by Glasgow Disability Alliance (Witcher, 2014) was set up by a user-led organisation to explore options for disabled people to enable increased choice and control in their day-to-day lives. The project focused on the core requirements of facilitating individual choice and independent living and found there was no obvious or consistent relationship between the amount of money that was spent on realizing individual choices and the scale of its impact. For example, some packages involved considerable investment for a British Sign Language interpreter to be available, whereas others required only a few hundred pounds for transport costs so that the individual could attend classes of their choice. Similarly, the test sites found no correlation between the size of the support package and the impact of SDS on individuals’ and families’ lives (Manthorpe et al., 2011). As Witcher (2014), along with Slasberg and Beresford (2015) argue, the success of SDS is in support that is truly person-centred and is clearly linked to self-defined outcomes. As such, the message of increased and more meaningful user-control from this type of initiative, underlines how control – or in the Norwegian concept of ‘citizen choices’ (Christensen and Pilling, 2014) should be a more central preoccupation in the implementation of SDS.

This links in with broader critiques by Beresford (2014) who argues that in England, commissioning authorities must ensure that the price paid in the ‘care market’ for services is enough to provide a personalised approach. In doing this, he emphasized the need for local areas to develop a network of providers who are able to deliver flexible and responsive services and not simply block contracts. Although the marketization of social care is arguably at a more advanced stage in England,
Scotland still has lessons to learn and securing a key role for user-led organisations would be an important way forward. In Norway, users are increasingly choosing a user-led organisation ‘Uloba’ as their service provider for personal assistance. As Christensen and Pilling (2014) show, these users are found to be significantly more satisfied with this model of support, than those who have chosen the municipality as the employer. However the promotion of this type of user-led model across the UK raises a note of caution in light of the wider challenges to sustaining activism as part of service provision and policy development amid long-term cuts to budgets. For Roy and Buchanan (2015), the climate of insecurity and uncertainty emanating from austerity has the potential to erode solidarity and trust, with the threat of unemployment making opportunities for service commissioners and providers to develop activism particularly difficult. Evidence from the initiatives documented by Beresford and Witcher above, as well as Roy and Buchanan’s own observations in recovery services show that this more negative outcome is not inevitable, but the austerity environment is undoubtedly more hostile to this vital support infrastructure.

In conclusion, at one level SDS is an uncomplicated idea that is impossible to disagree with, but as the experience of implementing SDS policy in Scotland has shown, there are ongoing challenges that need to be faced. To implement SDS in any meaningful way, requires a major paradigm shift in the relationship between users and professionals, and a refocus on the importance of the ethos of independent living. Further, the will to promote user centred practice and to embrace transformational change cannot be progressed in isolation from the austerity drive that is evident across Europe.
References


