The effectiveness of personal budgets for people with mental health problems: A systematic review

Short title: Personal budgets systematic review

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Abstract

Background: Personal budgets are a key policy priority in adult social care in England and are expected to become increasingly important in the care of adults with mental health problems.

Aims: This paper systematically reviews evidence for the effectiveness of personal budgets for people with mental health problems across diverse outcomes.

Method: The review, conducted in 2013, used the EPPI-Centre methodology for conducting a systematic review informed by Social Care Institute for Excellence guidelines. Data were extracted from studies and combined using meta-synthesis.

Results: 15 studies were included in the review which found mostly positive outcomes in terms of choice and control, quality of life, service use and cost-effectiveness. However, methodological limitations make these findings rather unreliable and insufficient to inform personal budgets policy and practice for mental health service users.

Conclusions: Further high quality studies are required to inform policy and practice for mental health service users which lags behind other adult social care groups in the use of personal budgets.

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Introduction

The aim of personal budgets is to put people at the centre of their own care and support. A personal budget recognises an individual’s strengths and preferences, and enables them to gain more choice in, and control over, the support they require to live their own lives. Helped in part by successful campaigning by disability rights groups and the Independent Living Movement, personal budgets have become a key health and social care policy in England (Department of Health, 2005, 2007; Her Majesty's Government, 2011).

Personal budgets have evolved in England since 2001 when Local Authority Adult Social Services Departments were required to offer eligible people cash direct payments to spend on care and support, which were frequently one-off payments for discrete items. ‘Individual budgets’ were subsequently developed, which were either a cash direct payment or a managed budget, or a combination of the two (Department of Health, 2005; Glendinning et al., 2008). These became known as personal budgets, which were rolled out throughout adult social care (Department of Health, 2007). However, widespread implementation has been accompanied by concerns that self-directed support (the process of support planning which elicits an individual’s goals and how to meet them), which underpins personal budgets, fails to live up to its ambitions (Slasberg, Beresford, & Schofield, 2012).

The use of personal budgets by people with mental health problems has been consistently lower than for other social care groups (Audit Commission, 2010; Davey et al., 2007; Riddell et al., 2005). For example, in 2011-12 only 14.6% of eligible mental health service users received a personal budget, compared to 58.8% of people with a learning disability, 47.9% with a physical disability and 45.2% of older people (Health and Social Care Information Centre, 2013). Organisational arrangements for the provision of mental health social care in England may have hampered the delivery of personal budgets to people with mental health problems (Larsen et al., 2013). Social care funding is means-tested and even where NHS and social care...
services are integrated, personal budgets require financial assessments and separate funding arrangements from NHS care (Audit Commission, 2010). Additionally, it has been proposed that there is a pervasive tension in mental health care between safeguarding against risk and the provision of user-directed care, which often involves some risk-taking (Carr, 2010). Consequently, mental health practitioners sometimes see people with enduring mental health problems as incapable of managing personal budgets (Carr, 2011; Taylor, 2008). Other barriers to implementation include the increased levels of bureaucracy that have accompanied personal budgets (Jacobs et al., 2013); concern amongst service users about managing personal budgets, and about the quality of support provided to help them with this (Newbronner et al., 2011; NHS Confederation, 2011).

Personal budgets may assist self-management programmes such as Wellness Recovery Action Plans (Copeland, 2002) by helping people to purchase services which support their recovery. Self-management is seen as increasingly important in long-term health conditions (Bodenheimer, 2003), and has become prominent in discourses of personal recovery in mental health services (Schrank, Bird, Rudnick, & Slade, 2012; Slade, 2009). Personal budgets also fit within current policy discussions about adult social care in England where co-production of services is becoming more common (Social Care Institute for Excellence, 2013). However, there are concerns that the marketplace for services and support to be purchased using personal budgets is under-developed which weakens their potential to work effectively (Spicker, 2013).

In spite of the low take-up of personal budgets to meet mental health social care needs, the Government have piloted personal health budgets for long-term conditions in England (Department of Health, 2009). Personal health budgets enhance choice and control over health care but, unlike social care personal budgets, are not means-tested. The evaluation of personal health budgets showed promising findings and they were particularly cost-effective for people with mental health problems (Forder et al., 2012).
Consequently, the Government has begun to introduce them into the NHS in England with plans to make them available in mental health services in 2015.

Despite the policy rhetoric, there is limited evidence about outcomes of personal budgets for people with mental health problems (The Health Foundation, 2010). Similar self-directed care systems such as ‘cash and counselling’ have improve satisfaction, service use and outcomes in the USA (Alakeson, 2010). However, Carr & Robbins (2009) reported that internationally there were few other schemes available for mental health service users. Although a prominent strand of government social care policy in England, there have been no systematic reviews of the literature examining the effectiveness of personal budgets for people with mental health problems. Therefore, this review aims to synthesise the evidence – qualitative and quantitative – about the outcomes of personal budgets for people with mental health problems.

**Method**

This review used the EPPI-Centre methodology for conducting a systematic review (Gough, Oliver, & Thomas, 2012), informed by the Social Care Institute for Excellence (SCIE) systematic review guidelines (Rutter, Francis, Coren, & Fisher, 2010). The development and conduct of the review was further informed by an Expert Advisory Group (EAG) with a range of research and clinical expertise, including service users and carers.

The search strategy was agreed by the research team in consultation with the EAG and refined by pilot searches using potential search terms. The final list of search terms used in the review were: cash and counselling; cash for care; consumer directed care; direct payment*; indicative allocation; individual budget*; individual service fund*; managed account*; managed budget*; notional budget*; personal budget*; personal health budget*; personalisation; personalised care; personalization; person centred;
pooled budget*; recovery budget*; resource allocation system (RAS); self-directed assessment*; self-directed care; self-directed support; support plan*; and virtual budget*. Terms were adapted according to the search functions and syntax of each database.

28 databases were searched; citation tracking and web searching was used to identify potentially relevant studies; selected social care journals and industry publications were hand-searched; and studies suggested by the EAG and other researchers/clinicians working in the area of personalisation were also sourced. The searches covered the full range of publication years up to April 2013.

Identified studies were screened by title, abstract and full-text and considered for inclusion in the review if they were published in English; included a clearly defined sample of adults with mental health problems aged 18-65 (irrespective of the presence of other disabilities); and presented ‘original’ empirical data/research, including qualitative studies of service users’ experiences of personal budgets. Studies were excluded if they were not available in English; included only people younger than 18 or older than 65 (as the use of personal budgets appears to be substantively different in these age groups and requires separate reviews); included only a sample of people with a disability where there was no evident mental health problem; focused only on person centred counselling or therapy (with no personal budget), or professional or organisational views; did not present separate results for people with a mental health problem; were opinion or comment pieces, or re-presented original research; or focused only on people’s experiences of the processes involved in delivering personal budgets with no outcome data presented.

To check inter-rater reliability of the application of the inclusion/exclusion criteria, the titles and abstracts of 20 randomly selected articles were double-screened by another researcher, as recommended by SCIE (Rutter et al., 2010). Additionally, the full text of ten studies was reviewed by three additional researchers. Discrepancies which arose were discussed and referred to the principal investigator and the EAG, and the
selection criteria were amended accordingly. Full-text studies which were not available to the researchers were obtained directly from the authors.

Data were extracted from the included studies using a standardised tool adapted from the EPPI-Centre Keywording Sheet (EPPI-Centre, 2002) and the SCIE Data Extraction Tool (Rutter et al., 2010). This tool facilitated the collection of details about the nature of the study, intervention and the outcomes. The tool also included a number of possible coding categories that was used to facilitate data analysis.

The quality of the included studies was appraised in order to assess the extent to which they had met the established norms for studies with those designs and reliably answered the question under investigation. Each tool included a number of domains which were considered to constitute a ‘low’, ‘moderate’ or ‘high’ risk of bias. There was also an overall risk of bias assigned to each study which comprised an average of the domain scores.

The Cochrane Collaboration’s risk of bias tool (Higgins et al., 2011) was adapted for use in the assessment of the randomised controlled trials and quasi-experimental studies. The Critical Appraisal Skills Programme (CASP) appraisal tool (National Collaborating Centre for Methods and Tools, 2011) was used in an abbreviated form to assess the qualitative studies, and Webber’s (2011) quality appraisal tool for cross-sectional surveys was also used. The latter tools were adapted to approximately map onto the categories of bias used in the Cochrane tool to allow for comparisons. Where studies used a mixed design, each part of the study was considered separately, such as the Personal Health Budgets evaluation (Davidson et al., 2012; Forder et al., 2012).
The variety of methodologies and analyses used in the studies eligible for review precluded the use of any meta-analysis (Harden & Thomas, 2005). Therefore, using the data extraction tool to form the basis of a narrative synthesis, findings were synthesised according to different outcome domains.

Results

15 studies met our inclusion criteria and the selection process is summarised in figure 1 below.
Figure 1: Application of selection criteria

Identification of potentially relevant studies through electronic databases (n = 27,531)

Additional studies identified through hand searching, citation tracking, Advisory Group input (n = 1641)

Duplicates screened and removed (n = 11,126)

Potential included studies after duplicates removed (n = 17,193)

Excluded at screen 1: titles and abstracts (n = 16,960)

Potential studies included in screen 2 (n = 233)

Excluded at screen 2: full text (n = 218)
  - No original data – eg. a review/commentary (n = 91)
  - No MH sample (n = 40)
  - Does not present results for a MH sample (n = 13)
  - Not an evaluation of personal budgets (n = 70)
  - Not about service user views (n = 5)

Studies included after screen 2 (n = 15)
Descriptions of the studies included in the review can be found in table 1 below. All studies were conducted in the UK or USA. Only two were RCTs, though an additional four used quasi-experimental methods. Only seven studies reported follow-up data. 1,135 people participated in 14 of the studies (range=2-412). The mean ages of samples ranged from 42–53 years and the proportion of people of black and minority ethnic (BME) origin ranged from 3%-50%. The proportion of females ranged from 26-70%. Participants were not defined by their sexual orientation. Studies evaluated individual budgets (n=2), recovery budgets (n=1), personal budgets (n=5), direct payments (n=1), personal health budgets (n=2) and ‘cash and counselling’ (n=4) as defined by their authors.
<table>
<thead>
<tr>
<th>Study No.</th>
<th>Study, year (country)</th>
<th>Study Design</th>
<th>Sample Size</th>
<th>[1]Age (2)Gender (3)BME</th>
<th>Outcome measures</th>
<th>Control Group</th>
<th>Intervention</th>
<th>[1]Length of follow-up (2)Length of budget (3)Payment used (4)Items purchased</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Glendinning et al, 2008 (UK)</td>
<td>RCT</td>
<td>n=131</td>
<td>(1)Mean=49 (2)46% female (3)5% BME</td>
<td>ASCOT (Netten, Forder, &amp; Shapiro, 2006); GHQ-12 (Goldberg &amp; Williams, 1988); Perceived QOL (Bowling, 1995); Self-rated health (based on Robine, Jagger, &amp; Romieu, 2003); Whether views have changed on what could be achieved in life; Overall satisfaction with support and specific aspects of quality based on User Experience Survey (Jones, Netten, Francis, &amp; Bebbington, 2007). Cost effectiveness measured with ASCOT and GHQ Care as usual</td>
<td>Individual Budgets</td>
<td>Care as usual</td>
<td>(1)6 months (2)Unclear (3) 71% DP; 26% LA (4)65% leisure, 57% social care; 31% PA</td>
</tr>
<tr>
<td>2</td>
<td>Homer et al, 2008 (UK)</td>
<td>Qualitative</td>
<td>n=2</td>
<td>Not given</td>
<td>Study-specific semi-structured interview schedule.</td>
<td>None</td>
<td>Individual Budgets</td>
<td>(1)None (2-4)Unclear</td>
</tr>
<tr>
<td>3</td>
<td>Coyle, 2009 (UK)</td>
<td>Qualitative</td>
<td>n=7 (n=4 at follow-up)</td>
<td>(1&amp;3)Not given (2)Female=2</td>
<td>Use of a 'narrative frame' which is not described in any detail.</td>
<td>None</td>
<td>Recovery Budgets</td>
<td>(1)7-8 months and 10-11 months (2&amp;3)Unclear (4)Most spent on IT equip, Gym, driving lessons, bike, course, home improvements</td>
</tr>
<tr>
<td>4</td>
<td>Cheshire West &amp; Chester Council, 2010 (UK)</td>
<td>Cross-sectional survey</td>
<td>Total sample size not given</td>
<td>Not given</td>
<td>Study-specific questionnaire</td>
<td>None</td>
<td>Personal Budgets</td>
<td>(1)None (2)Unclear (3)MH and LD had money paid into other’s a/c more than other groups (4)Unclear</td>
</tr>
<tr>
<td>5</td>
<td>Eost-Telling, 2010 (UK)</td>
<td>Qualitative</td>
<td>n=11</td>
<td>(1 &amp; 3) Not given (2)Female=3</td>
<td>The narrative frame or interview schedule is not clear.</td>
<td>None</td>
<td>Personal Budgets</td>
<td>(1)None (2)Unclear (3)Unclear (4)Total sample leisure</td>
</tr>
<tr>
<td></td>
<td>Hatton &amp; Waters, 2011 (UK)</td>
<td>Cross-sectional survey</td>
<td>n=78</td>
<td>Not given</td>
<td>POET(Hatton&amp;Waters, 2011)</td>
<td>None</td>
<td>Personal budgets</td>
<td>(1)None (2) 43% PB &lt;1 yr, 34% 1-3 yrs, 22% &gt;3 yrs (3)67% DP (4)Unclear</td>
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<tr>
<td>7</td>
<td>Lawson et al, 2010 (UK)</td>
<td>Cross-sectional survey</td>
<td>n=17</td>
<td>Not given</td>
<td>Short multiple-choice study-specific questionnaire</td>
<td>None</td>
<td>Personal Budgets</td>
<td>(1)None (2-4)Not clear</td>
</tr>
<tr>
<td>8</td>
<td>Rogers, 2009 (UK)</td>
<td>Qualitative</td>
<td>n=4 service users; n= 2 carers</td>
<td>(1&amp;3)Not given (2)Female (n=3 service users, n=1 carer)</td>
<td>Semi-structured interview tool</td>
<td>None</td>
<td>Personal Budgets</td>
<td>(1)None (2 - 4)Unclear</td>
</tr>
<tr>
<td>9</td>
<td>Spandler &amp; Vick, 2004 (UK)</td>
<td>Mixed: qualitative and cross-sectional survey</td>
<td>Qualitative (n=27); Questionnaire (n=16).</td>
<td>Not given</td>
<td>Semi-structured interview adapted from Witcher et al(2000)&amp;Ridley and Jones(2002) Quality of life questionnaire informed by the Lancashire QoLP(Oliver, 1991).</td>
<td>None</td>
<td>Direct Payments</td>
<td>(1)None (2)Mean=8.5 months (3)All DPs (4)PAs, then leisure, education, domestic support, pooled budget for arts group</td>
</tr>
<tr>
<td>10</td>
<td>Forder et al, 2012 (UK)</td>
<td>Quasi-experimental controlled trial</td>
<td>N=412 Intervention (n=228); Control (n=184)</td>
<td>(1)Intervention mean=45, control mean=53 (2)49% female (3)10% BME in intervention group, 3% in control</td>
<td>ASCOT;GHQ-12;Perceived QoL;Self-rated health(based on Robine et al., 2003); Whether views have changed on what could be achieved in life; EQ-5D(Brooks, Rabin, &amp; de Charro, 2003); Subjective well-being scale based on Dolan et al(2010)</td>
<td>Care as usual</td>
<td>Personal Health Budgets</td>
<td>(1)12 months (2)Unclear (3)27%DP, notional 8%, 3rd party – 63% (4)Social care, well-being, other health and therapy &amp; nursing</td>
</tr>
<tr>
<td>11</td>
<td>Davidson et al, 2012 (UK)</td>
<td>Qualitative</td>
<td>n=9 at 3 months, n=8 at 9</td>
<td>Not given</td>
<td>Semi-structured topic guide</td>
<td>None</td>
<td>Personal Health Budgets</td>
<td>(1)3&amp;9 months (2-4)Unclear</td>
</tr>
<tr>
<td>12</td>
<td>Shen et al, 2008 (USA)</td>
<td>RCT</td>
<td>n=228 Intervention (n=109) Control (n=119)</td>
<td>(1)70% aged 40-64 (2)70% female (3)50% White</td>
<td>No specific outcome measures were used, though the following outcomes were documented: the use of personal care services; perceptions of caregivers’ reliability and attitude; satisfaction with overall care arrangements; perceived unmet needs; adverse events; and satisfaction with life.</td>
<td>Care as usual</td>
<td>Cash and Counseling</td>
<td>(1)9 months (2)Unclear (3)Via fiscal intermediary (4)Only can be spent on personal assistance</td>
</tr>
<tr>
<td></td>
<td>Study Methodology</td>
<td>Study Details</td>
<td>Sample Size</td>
<td>Data Collection</td>
<td>Program Details</td>
<td>Key Outcomes</td>
<td>Key Findings</td>
<td></td>
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</table>
| 13 | Teague & Boaz, 2003 (USA) | Quasi-Experimental – with control | n=21 (Intervention: n=13; Control: n=8) | Not given | Structured interviews, questions taken from the Experiences of Care and Health Outcomes survey (Eisen et al., 1999), Working Alliance Inventory (Horvath & Greenberg, 1989), and others on service availability and use; perceptions of self-directed care; recovery goals; and outcomes. | Those who began the enrolment process but later declined | Based on Cash and Counseling | (1) None  
(2) Unclear  
(3) Fiscal intermediary  
(4) Mental health, physical health, housing, education/employment, enhancements (eg food, clothes, PA) |
| 14 | Cook et al, 2008 (USA) | Quasi-experimental (pre-post-design) | n=106 | (1) Mean=42.4  
(2) 61% female  
(3) 38% BME | Client outcome data compiled by the state mental health authority and GAF (Endicott, Spitzer, Fleiss, & Cohen, 1976) | None | Based on Cash and Counseling | (1) 1 year  
(2) 10.5 months  
(3) Fiscal intermediary  
(4) Traditional clinical services, clothes, food, non-traditional wellbeing services, uncovered medical expenses |
| 15 | Spaulding-Givens, 2011 (USA) | Quasi-experimental (no control) | n=80 | (1) Mean=51.4 ;  
(2) 61% female  
(3) 24% BME | Progress towards goal attainment; no of days in community in past 30 days; no of days worked in last 30 days, and income earned from that work. | None | Based on Cash and Counseling | (1) 1 year  
(2) 4.1 years  
(3) Fiscal intermediary  
(4) Transportation, housing/rent, utilities, dentist, medication, counselling, computers, clothing and food |

**Key to abbreviations used in table:**

BME=Black and minority ethnicity; DP=Direct payment; LD=Learning disability; MH=Mental health; LA=Local Authority; PA=Personal assistant; PHB=Personal health budget
None of the studies was considered to have a low risk of bias, with seven appraised as having a high risk of bias. A common limitation was a relatively small sample size, which impacted on analysis. Recruitment bias was evident in ten studies. The comparability of interventions in the multi-site studies (nos.1,2,6,9,10,11—see table 1) was made problematic by the different implementation of funding mechanisms at various stages of each study. None of the studies with quantitative data contained fully validated measures (for example, a perceived quality of life measure (Bowling, 1995) used in both Glendinning et al (2008) and Forder et al (2012) had not been validated for use with people with mental health problems) and none of the qualitative studies reflected on the impact that the researchers’ position potentially had on data collection or analysis. Finally, detail about analysis was generally lacking. The results of the studies are summarised in table 2.
## Table 2: Study findings

<table>
<thead>
<tr>
<th>Study, year (country)</th>
<th>Main outcome / Summary of findings</th>
<th>Limitations</th>
<th>OVERALL RISK OF BIAS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>INDIVIDUAL BUDGETS</strong></td>
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</tbody>
</table>
| Glendinning et al, 2008 (UK) | Intervention group reported significantly higher perceived quality of life (p<0.05)  
IBs appear more cost-effective | Relatively small sample; a number did not have budgets in place; significant over-representation of DP users; QoL measure not validated for MH; no benchmark nor acceptability curves for cost-effectiveness analysis; IBs implemented differently across sites; | MODERATE            |
| Homer et al, 2008 (UK, Scot) | Budgets enabled both participants to live a fuller life than otherwise | Very small sample, sampling strategy unclear; only included people in ‘critical need’; unclear method of analysis; budgets implemented differently in each area; | HIGH                |
| **RECOVERY BUDGETS**   |                                    |                                                                                                                                                                                                            |                     |
| Coyle 2009, 2011 (UK)  | Narratives before receiving budget were ‘regressive’, in contrast with ‘progressive’ post-budget narratives. Holding budget described as “entirely beneficial” and was key to success. Hope and recovery were core narratives. | No description of sampling method, narrative frame or researcher/interviewers’ stance; few negative/ambivalent themes; some of the items procured could have been obtained through other existing means; | MODERATE            |
| **PERSONAL BUDGETS**   |                                    |                                                                                                                                                                                                            |                     |
| Cheshire West & Chester Council, 2010 (UK) | Choice and control over life, and engagement in social and leisure opportunities had got worse, compared to other groups. Paid work had improved (of those in paid work) compared to other groups. | Sample and sampling strategy not described; no power calculations or discussion of response bias; no analysis described; no indication of significance or lack thereof of the results; only partial results presented. | HIGH                |
| Eost-Telling, 2010 (UK) | Narratives reported were all positive. Authority - more in control, confident; Direction - a new direction/motivation; Contribution - contributing more to society; Serendipity - positive knock-on effects in other areas of their lives | Sample recruited from one person’s case list; unclear about the interview frame, analysis, researcher’s stance – interviewer was member of clinical team; absence of negative or neutral narratives. | HIGH                |
| Hatton & Waters, 2011 (UK) | The following were reported as being better or a lot better: being in control of support, being independent (74%); in control of life (73%); mental wellbeing (70%); feeling safe (68%); relationships with family (64%); physical health (62%); relationships with friends (53%).  
Majority endorsed ‘no difference’ for getting and keeping a job, volunteering and community health, and in choosing where/with whom to live. | Over-representation of DP users; measure used not validated; the analysis and the statistics for group comparisons not given; generally results varied greatly across sites. | HIGH                |
<p>| Lawson et al 2010 (UK) | People with MH problems and LD were more likely to report improvements in their health and taking an active part in the community. | Small sample; MH over-represented in sample relative to the population norm; no information on questionnaire, data collection, nor the statistical analysis. | HIGH                |
| Rogers,                | High level of satisfaction when support plans in place. For some, cycle of | The sample size is very small and only half had budgets; unclear what the | HIGH                |</p>
<table>
<thead>
<tr>
<th>Year (Country)</th>
<th>Description</th>
<th>Analysis/Implications</th>
</tr>
</thead>
<tbody>
<tr>
<td>2009 (UK)</td>
<td>admissions broken, one moved out of residential care. One felt a good service had been withdrawn, and another felt they had never had an effective service and still not. Generally, once package in place, less contact with staff.</td>
<td>analysis process was or impact of researchers on it; overall conclusion is not apparently linked to the analysis.</td>
</tr>
<tr>
<td><strong>DIRECT PAYMENTS</strong></td>
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<tr>
<td>Spandler &amp; Vick, 2004 (UK)</td>
<td>Quality of Life questionnaire - positive impact on: self-worth (n=15); power and control (n=14); Independence (n=13); mental health (n=13); contact with friends/relatives (n=9). Only negative was one persons’ marital relationship. Qualitative – Increased autonomy, control, self-esteem; Increased social participation; Decrease in hospital admissions; Increased choice &amp; control – some found it hard to make choices, experienced greater uncertainty.</td>
<td>Small sample sizes; stable/trustworthy/articulate more likely offered DPs; length of time people had DPs in place not considered long; sites pre-agreed what DPs could be used for; decisions service driven; variations in implementation across sites; MODOERATE</td>
</tr>
<tr>
<td><strong>PERSONAL HEALTH BUDGETS</strong></td>
<td></td>
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</tr>
<tr>
<td>Forder et al, 2012 (UK)</td>
<td>Impact of receiving personal health budgets on outcomes: not significant - Cost analysis – ‘indirect costs’ were found to be significantly lower for PHB holders (at 5% level) - Cost-effectiveness – PHBs showed higher ASCOT –measured net benefits than conventional services (at 10% significance).</td>
<td>Relatively small sample size, low follow-up rate; PHB group less complex needs and were significantly younger than control; no cost-effectiveness threshold for ASCOT ;different sites implemented budgets differently; MODOERATE</td>
</tr>
<tr>
<td>Davidson et al, 2012 (UK)</td>
<td>Mostly improvements in well-being, stress, managing condition day-to-day, emergency service use and preparing for ill-health. A few negative responses also included mental health not improving as would like, no improvement, mental health had deteriorated.</td>
<td>Small sample, Recruitment bias as above; not all had PHB at 9 months; early stages report, may not be typical; no discussion of researcher impact on interview or analysis. Different sites implemented differently. MODOERATE</td>
</tr>
<tr>
<td><strong>CASH AND COUNSELING-BASED</strong></td>
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<tr>
<td>Shen et al, (2008) USA</td>
<td>More likely to be very satisfied with overall care arrangements (p&lt;0.001), and way life is being spent (p&lt;0.05). Less need for routine health care at follow-up (p&lt;0.01).</td>
<td>Number of people on budgets small; identifying sample from Medicaid claims problematic as people may not have made them in the previous year; unclear recruitment and outcome measures; proxies excluded. MODOERATE</td>
</tr>
<tr>
<td>Teague &amp; Boaz, 2003 (USA)</td>
<td>15% reported very important services were not available to them, compared to 87% of control group (p&lt;0.01), and were moving more quickly towards achieving goals (p=0.04). Also, increase in mental health (p=0.05) and independence (p=0.02).</td>
<td>Small sample; no well-defined comparison group; interview sample may not be representative; service participants seen as “higher functioning” than local MH population. MODOERATE</td>
</tr>
<tr>
<td>Cook et al, 2008b (USA)</td>
<td>More days in the community (than inpatient/forensic) compared to year before program (p&lt;.01). Significantly higher GAF Scale scores in the year after joining the program (p&lt;.001).</td>
<td>Methodology overall, outcome measures, follow-up periods for all measures, and whether all data has been reported are not clear. HIGH</td>
</tr>
<tr>
<td>Spaulding-Givens, 2011 (USA)</td>
<td>Disability Index (p=0.02) was significant –the more severely rated this score, the lower the amount of money spent.</td>
<td>Sample under-powered; input errors and missing data reduced this further; service users not representative of wider mental health population.; one measure was phased out during study; MODOERATE</td>
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</table>
We identified and synthesised four outcome domains:

(i) Choice and control of care and support

Four studies (nos.5,6,9,13) reported an increase in the levels of perceived choice and control experienced by people with mental health problems, including feeling more in control of life and support (nos.5,6,9), and feeling more confident, independent and powerful (nos.5,6,9, 13). However, one study (no.4) found that relative to other social care groups, mental health service users felt less in control of their care and support when in receipt of a personal budget. Two studies found evidence of an increase in choice, including flexibility with how time and resources were spent (no.9) and availability of services to budget holders (no.13). However, the increase in choice was accompanied by feelings of uncertainty in some service users (no.9), particularly when they found it difficult to articulate their needs.

(ii) Impact on life

Personalised care improved quality of life/overall satisfaction (nos.1,2,8,12); community participation (nos.5,7,8); physical health (nos.6,7,12); goal achievement (no.13); and greater sense of hope and recovery (nos.3,9). Three studies found benefits for mental health (nos.6,9,13) and one had mixed findings (no.11). Two studies reported generally better relationships with people (nos.6,9), though this was not the experience of all participants (no.9). Finally, one study (no.4) found that a personal budget helped people to keep paid work relative to other groups, but another (no.6) showed a large majority for whom it had no impact on employment.

(iii) Service Use

No study reported an increase in the use of inpatient services. As expected, five studies (nos.8,9,10,11,14) reported a decrease in community mental health service use.
Economic evaluations

Only two of the studies (nos.1,10) reported cost-effectiveness analysis. One found that personal health budgets were cost-effective (no.10), and the other that individual budgets appeared to be cost-neutral (no.1).

Discussion

This review provides some evidence that personal budgets can have positive outcomes for people with mental health problems, but a large number of methodological shortcomings limit the extent to which these findings can be interpreted. Further limits of interpretation come from the different payment and support mechanisms in the studies included in the review, and the different contexts in which they are applied.

The quality of the studies included in this review was moderate at best, with seven low quality studies, which is rather worrying given the significant investment in research and evaluation on personal budgets over the last ten years. Many personal budget evaluations had a high risk of bias, including those by the organisation charged with the delivery of self-directed support in the UK. The mental health sample within large studies included in this review was quite small, which impacted on the size of effect needed to demonstrate statistical significance, or artificially magnified others. Recruitment bias in these studies suggests that some of the participants were less ‘complex’, which causes further extrapolation problems. Also, the longest follow-up period in any of the studies was one year which, with difficulties in setting up personal budgets, led to many participants not being in receipt of their budgets at follow-up. Within some studies the funding mechanisms were deployed differently between sites. Outcome measures differed in their validity and reliability, making it difficult to compare studies and reach conclusions. Also, outcomes for people of BME origin, or lesbian and gay people, are largely unknown and deserve further exploration.
Our review screened over 17,000 studies against our inclusion criteria making it possible that potentially relevant studies were inadvertently excluded. US evaluations of brokerage were often unclear about funding mechanisms so it is possible that some which had personalised funding, but did not mention it, were screened out. Further, it was difficult to distinguish between process and outcomes in some studies. Evaluations of experiences of self-directed support, for example, were not included unless they referred to outcomes for personal budget recipients.

To the best of our knowledge, this is the only systematic review of the international literature on the effectiveness of personal budgets for people with mental health problems. It has found generally positive outcomes for mental health service users in terms of choice and control, impact on quality of life, service use, and cost-effectiveness. However, methodological limitations make these findings rather unreliable and insufficient to inform policy and practice. There is a need for large high quality experimental studies in this key policy area to inform personal budget policy and practice with people with mental health problems.

References


