The Health Benefits of Financial Inclusion: A Literature Review

Report for NHS Greater Glasgow and Clyde

Louise Dobbie and Morag Gillespie
Scottish Poverty Information Unit

May 2010
The Scottish Poverty Information Unit believes that poverty is caused by the unequal distribution of opportunities and resources rather than the lack of resources in society. SPIU aims to assist those committed to eradicating poverty in Scotland through robust policy analysis, quality research and widespread dissemination of poverty information. SPIU seeks to work in partnership with others towards the goal of reducing poverty and extending social justice in 21st Century Scotland.

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NHS Greater Glasgow and Clyde is committed to tackling the effects of structural inequalities. Its Equality Scheme ensures accountability for leadership and ownership for legislative requirements around this. Work across NHSGGC on employability and financial inclusion ensures that services intervene to ameliorate the effects of socio-economic inequality. The NHSGGC Financial Inclusion Group oversees this work and has a remit to develop the evidence base on financial inclusion, including the most effective interventions; establish connectivity between services; influence policy and strategy; develop performance indicators; create opportunities to share tools and developments that support financial inclusion and develop appropriate pathways between acute and community services.

Reference is made in this report to documents produced within NHS Greater Glasgow and Clyde. Further information about these and copies are available from:

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Report Conventions

**Advice Services:** This term is used to incorporate services that deliver welfare benefits or welfare rights advice (generally restricted to advice about welfare benefits and income maximisation work); debt or money advice (includes advice on a range of money related issues, including debt and income maximisation work) and general advice (including CAB services that provide advice in addition on issues such as employment, housing, taxation and utilities).

**Financial Exclusion** is concerned with “the lack of access to and use of a range of financial services”.

Exclusion can arise from a complex set of overlapping barriers and for a range of reasons including access, terms and conditions, marketing and price and self-exclusion. People who are financially excluded tend to be poor, live in relatively deprived areas and have certain characteristics such as being unemployed, sick, a disabled person, lone parent or single pensioner. As well as the greater likelihood of being poor, financial exclusion can mean people have lack a bank account or insurance, use high interest credit and pay more for utilities.

**Financial Inclusion:** This study uses the definition in the Scottish Executive Financial Inclusion Action Plan, 2005:

> “access for individuals to appropriate financial products and services. This includes people having the skills, knowledge and understanding to make best use of those products and services. Financial exclusion is often a symptom of poverty as well as a cause.”

**Abbreviations**

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>CAB</td>
<td>Citizens Advice Bureau</td>
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<td>CHCP</td>
<td>Community Health and Care Partnerships</td>
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<td>CPN</td>
<td>Community Psychiatric Nurse</td>
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<td>DRS</td>
<td>Development and Regeneration Services</td>
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<td>GAMH</td>
<td>Glasgow Association for Mental Health</td>
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<td>GAIN</td>
<td>Glasgow Advice and Information Network</td>
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<td>GCC</td>
<td>Glasgow City Council</td>
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<td>HADS</td>
<td>Hospital Anxiety and Depression Scale</td>
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<td>NHP</td>
<td>Nottingham Health Profile</td>
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<td>NHSGGC</td>
<td>NHS Greater Glasgow and Clyde</td>
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<td>RCT</td>
<td>Randomised Control Trial</td>
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<td>REA</td>
<td>Rapid Evidence Assessment</td>
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<td>SF 12/36</td>
<td>Short Form 12/36</td>
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Section 1: Background and Methods Used

Introduction

This report was commissioned by NHS Greater Glasgow and Clyde (NHSGGC) and produced by the Scottish Poverty Information Unit. The aim of this review is to provide a summary of evidence on the health benefits of financial inclusion and to assess the need for more research.

The links between poverty and ill-health are well established and people living with long term ill-health or disability are more likely to be living in poverty, a key factor in poorer health outcomes that have far-reaching effects on individuals and their families. However it has been argued that less is known about the pathways that link ill-health to poverty or how they may be broken. In particular, limited attention has been given to social participation, access to non-labour income, and other knock-on effects.4

The specific association between poor mental health and poverty is also recognised and there is a strong association between debt and poor mental health,5 with consequences for the relationship between mental health and financial inclusion. Cullen et al have highlighted that the anxiety and stress that accompany debt are known to impair health and affect relationships with family and friends.6 One quantitative research study highlighted that:

“The fact that a quarter of people with mental disorder were in debt has direct implications for effective clinical assessments and care planning, as well as for awareness in debt counselling agencies, utility companies and financial organizations.”7

Current evidence shows that the health inequality gap is widening - income inequality has risen in each of the last three years and is now at its highest level since our comparable time series began in 1961.8 The recent economic downturn is likely to worsen the situation for the most deprived communities and excluded groups including women, black and minority ethnic people, disabled people, homeless people, refugees and asylum seekers. The situation of children in families vulnerable to unemployment and other disadvantage is also likely to be affected. There is now greater acknowledgement of the potential for measures that tackle financial exclusion to reduce health inequality and tackle the social determinants of ill-health. A key recommendation in the Equally Well framework stated:

“Universal public services should build on examples of effective financial inclusion activity, to engage people at risk of poverty with the financial advice and services they need. Removing the stress caused by debt will improve people’s health and wellbeing.”9

Welfare benefits in the UK are complex and potential beneficiaries may face a range of barriers to accessing entitlements to benefits, including, for example, age, health, transport, access, literacy or cognitive difficulties.10 Improving the take up of benefits entitlements, or income maximisation, has been a key area of investment by the NHS as one means of helping to address low income and poverty and contributing to
reducing health inequalities. This includes support for provision of services from within healthcare settings, particularly primary care.11

The development of the broader approach involved in financial inclusion is relatively recent and it embraces a range of services or activities including money advice and income maximisation work, awareness raising and service provision around, for example, savings or low cost credit, and financial capability or money management support and guidance. While advice is integral to financial inclusion, more holistic approaches are now recognised as essential.

However, in relation to health improvement, advice has been important to debate so far and is the most researched area of financial inclusion. It remains the most common financial inclusion activity located in health services and the number of projects and services delivered in health settings has grown. Most often evaluations and research on such projects and services have recorded activities and, for example, the types and amounts of benefits gained. However, most have not measured the health benefits for clients or the benefits to health services, including some research within the NHSGGC area.12, 13, 14

NHS Greater Glasgow and Clyde objectives on financial inclusion

The NHS has contact with people as part of their rehabilitation and self-care pathway and, therefore, an opportunity to support people in accessing wider social needs, such as welfare benefits entitlements. Research has identified that:

“Roughly two thirds of the population consult their general practitioners at least once a year. General practitioners and community nurses are exceptionally well placed to detect those who are suffering genuine financial hardship but they are not well equipped to give advice about the complex system of state social security benefits.”15

NHSGGC is committed to tackling the effects of structural inequalities. Its Equality Scheme ensures accountability for leadership and ownership for legislative requirements around this. Work across NHSGGC on employability and financial inclusion ensures that services intervene to ameliorate the effects of socio-economic inequality. The NHSGGC Financial Inclusion Group oversees this work and has a remit to develop the evidence base on financial inclusion, including the most effective interventions; establish connectivity between services; influence policy and strategy; develop performance indicators; create opportunities to share tools and developments that support financial inclusion and develop appropriate pathways between acute and community services.

To develop an inequalities sensitive health service NHSGGC wishes to skill health practitioners to understand the social issues and structural inequalities facing their patients, support patients with these and have the capacity to refer them to appropriate services. NHSGGC has involvement in partnerships that have access to a range of resources and staff who have knowledge of financial inclusion and can work together to join up the pathway for health and social care clients to improve their health outcomes.
Various parts of NHSGGC have piloted approaches to financial inclusion. In order to take a preventative, upstream approach to tackling these issues, the lessons from current practice need to be mainstreamed in a sustainable way. NHSGCC Financial Inclusion Group is coordinating a proactive approach and has developed an action plan, which it reviews regularly. A recent mapping exercise of current financial inclusion provision across NHSGGC found different levels of service provision and no consistent referral pathways but concluded that “there is a wealth of activity developing on financial inclusion in health settings which could be replicated or used to develop more consistent models”. The Financial Inclusion Group is developing service standards that will facilitate monitoring in the longer term across NHSGGC.

**Research Methods Used**

The broad aim of the study was to summarise the health benefits of financial inclusion and to assess the need for more research and it involved the following:

- A review of evidence of the health and quality of life impacts of financial inclusion initiatives, with particular reference to collating evidence of NHS-based interventions & the health benefits of these
- Exploring models of practice and learning to improve practice and identify evidence of the tools and barriers that exist
- Reviewing research methods used in existing studies
- Development of recommendations about future policy, practice and research

The research team undertook a Rapid Evidence Assessment (REA). REAs provide a balanced assessment of what is already known about a policy or practice issue. Systematic review methods are used to search and critically appraise existing research. They aim to be rigorous and explicit in method and so systematic. In comparison, REAs make concessions to the breadth or depth of the process by limiting particular aspects of the systematic review process. The method is appropriate for this review because the policy area is relatively new and there are few existing studies. Full details of the search strategy and analyses conducted are provided in Appendix 1. The REA included peer-reviewed journals and local monitoring and evaluation reports. It also involved review of other important sources such as government, in-house and commissioned reports, independent (non-journal published) research (e.g. studies commissioned by the Joseph Rowntree Foundation), and analyses or commentaries from stakeholders (e.g. ‘poverty lobby’ organisations, consumer groups). It focused on English language sources and data, and on reports published in the last 10 years, with an emphasis on UK studies. Evidence of the health impacts of financial inclusion was recorded where identified.

The data searches covered the time period from 1999 to 2009. This reflects the pace of change in the area of financial inclusion policy, which makes many older sources redundant. The review searched for material using the most important bibliographic social science databases and search engines, as well as following references and citations in the texts consulted.
Section 2: The Health Impacts of Financial Inclusion

This section reviews the information available on the impact of financial inclusion work on the health of individuals. It reflects the fact that most research relates to advice provision that addresses either welfare benefits or more general advice needs, including money advice. However, where it has been found, information is identified that relates to wider financial inclusion initiatives and their relevance for health services and improving health. The report also seeks to broaden discussion of the impacts of advice to embrace not only the physical and mental health outcomes of interventions for participants but also the wider effects on quality of life and wellbeing.

The links between poverty and poor physical and mental health are clear and improvements to health, wellbeing and quality of life are amongst key reasons for developing advice services linked to primary care. However, the causal pathways are complex and the way in which an increase in individual income can improve individual health is not clear. Referring in 2002 to a small number of studies that considered the health benefits of advice, Abbot argued that welfare benefits advice located in primary care can be expected to have limited impact as a health intervention. Rather, it offers more potential for relief from psychological stress for service users and may also make a small contribution to the physical health of individuals and to the social capital of communities.

Existing literature and evidence reviews

Since 2002, there has been a small number of reviews of the health impacts of advice. One review which included 55 studies was conducted by Adams et al in 2006. It focused on the health, social and financial impacts of welfare rights advice in healthcare settings. They identified that, in quantitative studies, the instruments used to measure health impacts included non-specific measures of general health such as the Short Form 36 (SF-36, a general health scale), the Hospital Anxiety and Depression Scale (HADS, a questionnaire commonly used to screen for anxiety or depression), the Measure Yourself Medical Outcome Profile scale (MYMOP, a patient generated wellbeing scale), the Nottingham Health Profile (NHP, a quality of life scale), and the Edinburgh Post-natal Depression Scale. Key points concerning quantitative studies relating to health and social outcome highlighted by the authors included:

- Amongst seven studies that had a comparison or control group - 11 of 72 separate comparisons reported were “statistically significant at the 5% level including comparisons relating to SF36 vitality, SF36 mental health, SF36 bodily pain, SF36 role functioning emotional (and) NHP emotional reactions”.

- Of six studies reporting non-financial results using recognised measurements scales and before and after study design, 6 of 59 separate statistical comparisons reported across 3 studies were found to be significant – SF36 vitality, SF36 role functioning emotional, SF36 mental health, SF36 general health, NHP pain and NHP emotional reactions.
Their assessment was that overall: sample sizes were small; follow up was limited to 12 months at most; and few statistically significant differences were found and these tended to relate to measures of psychological or social, rather than physical health.

Of the other studies identified, the authors saw little merit in seven which used in-house questionnaires with “little evidence of validation”. These “found consistently high levels of clients agreeing with statements concerning the positive impact of the advice on their health, quality of life and living situations”.18

In the 14 studies reporting qualitative data, most data were gathered through interviews with individuals including those receiving advice, advice givers and primary care staff. The most common use of qualitative data was to assess perceptions of the impact of advice and these showed:

- Advice based in healthcare settings, particularly primary care, was valued and it legitimised advice, improving access and reducing stigma
- Advice and benefits helped elderly clients maintain independence
- Advice decreased worry and anxiety and improved mental health and quality of life irrespective of whether individuals had an income increase as a result of advice
- Advice and increased benefits increased physical health and reduced use of health services.
- Advice was seen as ‘expert’ and professional
- Negative comments concerned GPs’ lack of awareness of the service, concerns about losing benefits and benefits increases being offset by deterioration of health due to long term illnesses

Adams et al described the evaluations found as having limited scientific rigour and linked this to scarce resources and skills. They concluded that there was a “theoretical potential” for advice to improve health, but they recommend more robust methodologies including longer follow-up, since the health and social effects of increased financial resources may take years to become apparent. They also advocated the use of randomised control trials. This issue is discussed further below (page 17).

Other reviews were identified which considered the health impacts of advice. In 2002, Greasley and Small19 considered the same evaluations of the benefits of welfare advice on the health and well-being of participants as were included in the more recent review by Adams et al. Wiggan and Talbot’s 2006 review of the impact of welfare rights advice included some consideration of the health impacts of advice. Their assessment was based more on a social model than Adams et al. While they discussed some of the same studies, they concluded that the literature indicated “significant improvements in mental health of clients following successful welfare rights
intervention and improvement in physical health, although these may be more limited".\textsuperscript{20}

A review of social support services for people affected by cancer, focusing on financial advice and support, highlights problems, bureaucracy and delays but identified less research that discusses the impacts of advice.\textsuperscript{21} However the study highlights recent positive developments in the MacMillan Cancer Care Advice project. Most recently, Green reviewed money advice giving methods in 2009 and considered their relative costs and effectiveness for different client groups. Research gaps she identified included: the impact of individual money advice approaches, understanding of the impact of debt advice on different population subgroups and debt advice outcomes in Scotland, but she did not discuss health impacts of advice.\textsuperscript{22}

**Individual Studies - research and evaluations**

This review has identified 16 key studies or journal articles which have reported health and social outcomes of financial inclusion interventions and these are summarised in Table 1. We have focused on studies published since the review conducted by Adams et al. However, a small number have been included here because they contain information relevant for the present review, which is concerned with financial inclusion activity, rather than a singular focus on the provision of advice in healthcare settings. These studies are annotated in the Table.

All of the studies identified discuss health impacts of advice provision. Two studies evaluated other approaches in addition, such as awareness raising sessions, money management guidance, or development activities.\textsuperscript{10,23} The studies generally focused on the provision of welfare benefits advice, including income maximisation work. However, Citizens Advice Bureaux (CAB) and many other advice services advise clients on a range of social, legal and welfare rights issues (including, for example, housing, employment, taxation and debt) and several studies highlight the wide range of issues addressed and the fact that individual clients may raise more than one problem.

The interventions involved: delivery of advice located in primary care services (seven studies); one project, discussed in two separate articles was located in a social services department;\textsuperscript{24,25} two were located within the advice providers premises; and one longitudinal study considered the health impacts of advice in both the mainstream CAB service and outreach in primary care.\textsuperscript{26} The remainder involved a mix of delivery methods including telephone advice, home visits following screening or referral by primary care and other staff. Half the studies had no specified target groups or aimed to reach a range of different service user groups, some others targeted people aged 60 or more and individual studies were concerned with people with chronic conditions, stroke patients and people with mental health problems.

The providers of advice were most commonly a mix of services, CAB services or local authority welfare rights, while one study involved a voluntary sector mental health service. The providers in two studies were unclear. Excluding the project monitoring data gathered by advice providers, half of the studies used qualitative methods and the remainder were evenly split between mixed method and
quantitative studies. Within these, the qualitative and quantitative elements of two studies were reported in separate articles and considered separately here.

In the studies with a quantitative element, the Short Form 36 (SF36) was used in five cases to measure health impact. However, more than one measure was used in some studies and the Hospital Anxiety and Depression Scale (HADS) was used in two studies, while a further six instruments were each identified once.

Sample sizes varied considerably, including attrition rates in those studies that had follow-on interviews. Only two studies used randomized control trials (RCTs), one of which was abandoned because of sample attrition. Sample sizes also varied considerably in the qualitative studies. Most involved interviews with individuals accessing services, although in four studies other groups included were staff from the advice services, primary care or other service providers.

The main message from across the studies is that both qualitative and quantitative methods identify benefits from advice in terms of improved mental health, reduced stress or anxiety and better quality of life, but there is less evidence of improvements to physical health. Relatively short follow-on study periods and other methodological issues are suggested to have contributed to modest results in some studies.

While Abbott et al identified that the benefits of increased income were modest but valuable, other studies stressed that increased income could help manage health better, improve lifestyle and help relationships, for example with children. One qualitative study identified that stroke patients' rehabilitation and speed of recovery was improved. Conversely, Moffatt and Scambler found that the poor health of the people they interviewed was attributed to factors that were considered "immune to the effects of money", except to the extent that more money affords greater capacity to manage health conditions.

The links between income increases and positive health changes are very complex. This raises questions about how important the nature of the health condition is in relation to the point of intervention in the patient pathway. For example, health may be affected more by income increases in acute health conditions or at the initial stages of diagnosis or falling ill (such as stroke or cancer) compared with more chronic health conditions for which income over the long-term or income inequality may be more important.

A further consideration is that advice and income maximisation may help to slow or prevent further deterioration in health. The links between poor mental health and other physical health conditions and the evidence of the positive impacts of advice on mental health from these studies, provide positive arguments for advice. However, the impact on physical health may need much longer term studies than have been possible thus far. In the short-term though, "helping to ease the problems of those who are both deprived and very unwell, even an imprecise intervention can be of value".

Campbell et al suggested that advice or other external factors may have played some part in health stability, as opposed to the increase in financial income per se. In one study “many participants spoke at length about having the chance to talk, and
having someone take an interest, suggesting that the welfare consultation itself had a therapeutic effect." The importance of access to ‘talking therapies’ was recognised in the design of the Health Plus Project in Bradford. In addition to providing advice, this project employed 2 full time equivalent counsellors to provide brief intervention therapy to patients suffering from mild to moderate mental health issues. However to better meet needs and demand for the service, the design was revised to include a larger multidisciplinary therapeutic team of 9 full-time posts with a team leader. In a similar vein, research in Scotland found that, for people seeking advice about debt, the impact of advice in reducing the stress and anxiety surrounding indebtedness can be important. This need not necessarily involve an increase in income, so much as helping people to make their financial situation manageable through a range of strategies from negotiating repayments through to bankruptcy.

**Broader Financial Inclusion Interventions**

The body of work identified has an almost exclusive focus on welfare rights advice and there are few studies referring to financial inclusion initiatives other than welfare benefits advice. For example within NHSGGC, the Health Improvement Team within Acute Planning developed a successful bid, following consultation, to fund a pilot project that would meet the money advice and financial inclusion needs of people affected by stroke. The pilot service was designed to meet the immediate financial needs of people affected by stroke within the acute setting and facilitate a transition to specialised community based financial inclusion services when and if required. The evaluation report highlighted that “feedback from all stakeholders noted a positive impact on patient rehabilitation and speed of recovery. In particular, NHS staff noted the direct impact financial worries could have on individuals affected by stroke and the crucial role they can play in helping people access support”. Unfortunately, further research is needed to support this finding and ascertain other element of the work which were not reported on, including the financial capability work.

Liddell conducted a review which considered recent evidence on tackling fuel poverty and the impacts on human health. One of the studies included was an evaluation of the Scottish Central Heating Programme which found limited impacts on health. Importantly, the evaluation found that the programme was not well targeted at low income groups since “as many as 50% of homes targeted by CHP were not fuel poor even though they passed all eligibility criteria”. Nonetheless, Liddell concluded on the basis of other existing evidence that such initiatives do have the potential to reduce the effects of poverty and contribute to reducing health inequalities:

> “The health impacts of tackling fuel poverty appear noteworthy, especially when taking into account that they derive from a source as seemingly tangential to human health as home heating and cavity wall insulation. Improved mental wellbeing amongst the elderly, when taken in conjunction with impacts on adolescents, children and infants that have the potential for a life-long legacy, provide an evidence base that supports the rationale for fuel poverty strategies of the past, present, and future.”
The wider literature on fuel poverty has not been reviewed here. However, people living in poverty face higher fuel costs than average, particularly if they are using prepayment or card meters and if they are repaying arrears. The options to help reduce fuel costs are, therefore, related and potentially important areas of activity to reduce financial exclusion, reduce costs and improve health.
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<th>Authors/Yr</th>
<th>Project</th>
<th>Methods</th>
<th>Health Impact Findings</th>
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| *Greasley (2003)*<sup>34</sup> | Evaluation of the Health Plus Project (6 advice workers) in 30 General Practices in Bradford. | Methods used included SF-36 and the Hospital Anxiety and Depression Scale (HADS) and an in-house questionnaire; focus groups with advice and health care staff; questionnaire to practice managers (132 at baseline, 64 at 6 months and 32 at 12 months 6) | SF36 and HADS) suggest welfare advice may improve health and quality of life for patients. Information from an additional questionnaire supported this, for example:  
  • 81% of patients who responded at 6 months (52/64) said the advice had helped them  
  • 21 of them received extra income; 15 of whom said that this had affected their health and how they feel – reducing anxiety/worry/stress about their financial situation and improving their quality of life  
  The project was effective at reaching people with mental health problems and older south Asian women - two thirds of service users were from minority ethnic groups |
| *Moffatt et al (2004)*<sup>37</sup> | Study of advice sessions in GP practices. Includes discussion of impact on health/ outcome measures | Qualitative study with CAB and 3 general practices, 14 semi-structured interviews, including 11 who gained increased income, age range 46-76 (research in 1997-8) | Everyone reported improvements or stability in mental health as a result of welfare advice.  
  The authors hypothesised that increased resources and the resulting increases in choice and control have three main effects: reducing psychological stress, which may be expressed in a range of ways, including decreased worry, pressure, anxiety and depression |
| *Abbott et al (2005)*<sup>28</sup> | Reports findings of a study designed to measure changes in individual health associated with income increase as a result of advice | Study 2000-01 in seven sites across England. Longitudinal observational study of advice to service users, comparing the health of those whose income increased with that of those whose income did not, using the SF-36 as an outcome measure. 345 people were interviewed at base line, 245 after 6 months and 201 after 12 months. They were generally in the second half of life, with one or more chronic conditions (74% = 45+, 26% = 15-44yrs) | The income increase group had significantly better outcomes in mental health and emotional role functioning at 12 months than those with no income increase. There were no other significant differences between groups at 12 months, and none at 6 months.  
  Over half of the income increase group said they were better able to pay bills, nearly half said they used transport more. Over a third said that they could now eat more and/or better food.  
  Argues that the impact on health of increased welfare benefits is modest, at least for people in the second half of life with chronic illness and/or disability, who tend to use primary care based services. However, the benefits of such small improvements in quality of life of deprived patients with chronic ill health should not be underestimated: qualitative evidence supports the value of such improvements to patients themselves - welfare benefits advice has a part to play in the holistic approach to the care of such patients. |
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<th>Author(s)</th>
<th>Methodology</th>
<th>Findings</th>
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<tr>
<td>Abbott and Hobby (2005)</td>
<td>Based on Abbott et al (2005). Explores associations between changes in health status and increased income from advice. Reports six cases in detail from larger study in 2000-1 to illustrate interplay between financial, social and health aspects for deprived patients. Structured interviews with advice recipients after the advice and 6 &amp; 12 months later.</td>
<td>Two patients recognised the role of an income increase in improving their health-related quality of life; in two cases, social and health issues outweighed the benefits of income increase; and two patients had, despite advice, experienced a reduction in income that they believe affected their health for the worse. Argue there is a clear if modest role for benefits advice as part of a holistic approach to health in primary care.</td>
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<td>Caiels and Thurston (2005)</td>
<td>Research to establish how successfully a GP outreach project had met its original objectives Mixed methods: Routine monitoring data (CAB 333 referrals); Pre- and post-service intervention questionnaire (96); Health status measurement (SF-12 - 90 pre- and 84 post-service intervention) Semi-structured interviews (6 service users, 7 primary health care professionals, 3 CAB advisers)</td>
<td>78% reported feeling less anxious after seeing the CAB adviser, 2% felt more anxious and 20% felt the same. Service users found the service accessible and advice was advantageous. Not able to show improvements in health as measured by the SF-12. However, contributed to gains in health-related quality of life, including increased income, reductions in stress and improved housing situation, factors relevant to addressing health inequalities and improving health-related quality of life. Advisers reported observing ‘real’ improvements in clients as a result of accessing the service, but were realistic about improving health among those with severe mental illness or disability.</td>
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<td>Mackintosh et al (2006)</td>
<td>Reports on impact of welfare rights advice over 24 months in primary care, discusses methods, including potential outcome measures. Single blind, pilot RCT in 2003 (126 aged 60+, 109 at follow-up). Control group had intervention after 6 month delay. Measurement: SF36, HADS, activity limiting long term illness, symptoms inventory, the Pittsburgh Sleep Quality Index and BMI measurement</td>
<td>There was little evidence of differences in health outcome measures between groups or over time. (may be due to study design) Possible reasons given include: the small sample size; inadequate lengths of time for additional welfare benefits to have health and psycho-social effects; many participants did not qualify for any benefits; and the outcome measures used may not have been the most appropriate: ‘these factors combined would have reduced the observed strength of any possible effects.”</td>
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<td>Moffat et al, (2006)</td>
<td>Qualitative study of the views of older people about the impact of welfare rights advice (study concurrent with Mackintosh et al, 2006) 25 initial interviews (2003) and 22 follow-up interviews after 18 months (2005) - schedules semi-structured and covered: changes in circumstances as a result of advice; perceived impact of material and/or financial benefits on mental and/or physical health; health-related behaviours; social benefits; and views about the associations between material resources and health.</td>
<td>4 categories of impact identified from additional financial resources : 1) raised ability to buy necessities 2) more able to afford occasional expenses 3) capacity to cope with future crises 4)more peace of mind Many mentioned that they had a healthier diet (able to afford better quality foods and not always cheapest) Findings included: Extra money or resources has no effect on an individual’s health. Poor health was attributed to factors considered to be immune to the effects of money. However, although being more able, for example, to afford daily necessities, keep up with friends and family did not directly alter health, it increased capacity to cope with ill health.</td>
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Caiels and Thurston (2005)  | Research to establish how successfully a GP outreach project had met its original objectives Mixed methods: Routine monitoring data (CAB 333 referrals); Pre- and post-service intervention questionnaire (96); Health status measurement (SF-12 - 90 pre- and 84 post-service intervention) Semi-structured interviews (6 service users, 7 primary health care professionals, 3 CAB advisers) | 78% reported feeling less anxious after seeing the CAB adviser, 2% felt more anxious and 20% felt the same. Service users found the service accessible and advice was advantageous. Not able to show improvements in health as measured by the SF-12. However, contributed to gains in health-related quality of life, including increased income, reductions in stress and improved housing situation, factors relevant to addressing health inequalities and improving health-related quality of life. Advisers reported observing ‘real’ improvements in clients as a result of accessing the service, but were realistic about improving health among those with severe mental illness or disability. |

Mackintosh et al (2006) | Reports on impact of welfare rights advice over 24 months in primary care, discusses methods, including potential outcome measures. Single blind, pilot RCT in 2003 (126 aged 60+, 109 at follow-up). Control group had intervention after 6 month delay. Measurement: SF36, HADS, activity limiting long term illness, symptoms inventory, the Pittsburgh Sleep Quality Index and BMI measurement | There was little evidence of differences in health outcome measures between groups or over time. (may be due to study design) Possible reasons given include: the small sample size; inadequate lengths of time for additional welfare benefits to have health and psycho-social effects; many participants did not qualify for any benefits; and the outcome measures used may not have been the most appropriate: ‘these factors combined would have reduced the observed strength of any possible effects.” |

Moffat et al, (2006) and Moffatt and Scambler, (2008) | Qualitative study of the views of older people about the impact of welfare rights advice (study concurrent with Mackintosh et al, 2006) 25 initial interviews (2003) and 22 follow-up interviews after 18 months (2005) - schedules semi-structured and covered: changes in circumstances as a result of advice; perceived impact of material and/or financial benefits on mental and/or physical health; health-related behaviours; social benefits; and views about the associations between material resources and health. | 4 categories of impact identified from additional financial resources : 1) raised ability to buy necessities 2) more able to afford occasional expenses 3) capacity to cope with future crises 4)more peace of mind Many mentioned that they had a healthier diet (able to afford better quality foods and not always cheapest) Findings included: Extra money or resources has no effect on an individual’s health. Poor health was attributed to factors considered to be immune to the effects of money. However, although being more able, for example, to afford daily necessities, keep up with friends and family did not directly alter health, it increased capacity to cope with ill health. |
<table>
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<tr>
<th><strong>Pleasance et al (2006)</strong>&lt;sup&gt;27&lt;/sup&gt;</th>
<th>Research project to establish impact of debt advice on people’s lives (including impact on health). Summarises results of 4 studies from the Impact of Debt Advice Research Project</th>
<th><strong>Study 1</strong>: quantitative interviews (176 clients at time of advice and 6 and 12 months later. Only 61 remained in the study at 12 months  <strong>Study 2</strong>: analysis of quantitative data from the 2004 English and Welsh Civil and Social Justice Survey (CSJS)  <strong>Study 3</strong>: 42 qualitative interviews with people with debt problems.  <strong>Study 4</strong>: RCT to explore whether debt advice would improve their life circumstances, relative to a similar ‘control’ group referred for telephone advice</th>
<th>Evidence was not conclusive but indicated levels of anxiety, general health, relationships and housing stability benefited from advice. <strong>Agency study</strong>: following advice, clients generally reported improvements in their health. At both six and twelve-month follow-up, about two thirds of clients said their health had improved a little or a lot since the time of initial interview. 75% at 6 months and 90% at 12 month follow-up said they felt health improvements stemmed from receipt of advice, but 11% reported changes for the worse 12months after getting advice. <strong>State Trait Anxiety Index</strong>: Self-reported general level of health also improved among advice agency clients between the time of initial advice and six and 12 month follow-up. Also saw small positive change in health status, as recorded through the five main EuroQol EQ-5D general health index questions. RCT not completed due to levels of attrition</th>
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<tr>
<td><strong>Campbell et al (2007)</strong>&lt;sup&gt;24&lt;/sup&gt;</td>
<td>Longitudinal postal survey of individuals aged 60 and over referred for specialist welfare benefits advice within social services and who were followed up at 5 months (over 2003–2004). Outcome measures included SF-36, the General Health Questionnaire-12 and the Barthel Index (postal version), along with questions relating to chronic illness 77 baseline questionnaires returned. 52 of the 69 (75%) remaining participants completed follow-up questionnaires. See Winder et al (2008) on associated qualitative study</td>
<td>Following advice, physical health (SF-36 and Postal Barthel) scores remained stable, with marginal improvement in the Role Limitation (Physical) SF-36 dimension and the single Change in Health question. An overall reduction in GHQ-12 scores, suggested an improvement in psychological wellbeing. Results were not affected by whether or not individual’s benefits increased. 44% respondents previously had benefits advice and many were already receiving benefits prior to assessment. Newly referred respondents showed improvements in the SF-36 dimensions of Social Functioning and the Change in Health single item, despite a smaller increase in financial benefits compared with those who were already known to the service.</td>
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<td><strong>Gillespie et al (2007)</strong>&lt;sup&gt;10&lt;/sup&gt;</td>
<td>Evaluation of 11 Money Advice for Vulnerable Groups pilot projects across Scotland</td>
<td>Methods used included: project data analysis, an advice quality check, focus groups with staff and managers, interviews with 111 service users and 42 with staff in other services; and follow up interviews with 37 service users and 13 staff. Not in health setting and no specific evaluation of health impacts</td>
<td>46% of follow-up interviewees said accessing money advice and being provided with appropriate support had improved their mental health and well-being. Improvements cited most often were reduced stress or feelings of anxiety and 16.7% said dealing with their financial worries had also improved their relationships. Argues that money advice can reduce ‘risk factors’ and strengthen ‘protective factors’ for positive mental well-being and address key principles of effective intervention including reducing anxiety, enhancing control, facilitating participation and promoting social inclusion.</td>
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<tr>
<td>Author and Year</td>
<td>Study Description</td>
<td>Methodology</td>
<td>Findings</td>
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<td>Turley and White (2007)</td>
<td>Aimed to assess role and value of advice for people with a debt problem</td>
<td>42 face to face depth interviews recruited from 82 participants who reported a money problem in the 2004 English and Welsh Civil and Social Justice Survey. Component of Pleasence et al (2006) study</td>
<td>Respondents described a range of effects from feeling stressed to experiencing more serious mental health difficulties, so having an improved emotional outlook as a result of receiving advice was invaluable to these respondents. Respondents described feeling reassured, less anxious and less depressed.</td>
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<td>NHSGGC (2008)</td>
<td>Evaluation of pilot financial inclusion service to meet immediate financial needs of people affected by stroke (acute setting) and facilitate a transition to community based services</td>
<td>Qualitative interviews with 42 patients and carers and unknown number of advice workers. Questionnaire responses from 12 NHS staff. Advice at each stage, so early intervention. Service provided transition support and continuity of adviser on discharge and holistic support covering all aspects of money advice</td>
<td>In total 93 patients and carers used the money advice service. This represented 21% of the stroke acute population within the 6 month pilot period. Of the 10 users who knew where to access support locally only one said they would access it. Reasons for not using it included not feeling well enough to get to the service, having communication difficulties, long waiting lists and embarrassment at having to go and ask. Patients and carers felt that the resulting stress and anxiety caused by money concerns had a negative effect on the speed of their rehabilitation. Feedback from all stakeholders noted a positive impact on patient rehabilitation and speed of recovery.</td>
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<td>Winder et al (2008)</td>
<td>Explore experiences of benefit advice from a specialist, social services-based team for older people and their carers and its effects on health/well-being. Qualitative study run in parallel to an associated longitudinal observational study examining the impact of welfare benefits advice on client health and well-being (Campbell et al, 2007) 27 interviewees people aged 60+ and carers</td>
<td>Interviewees described difficult life circumstances around the time of referral. The complexities of accessing advice and of understanding the benefits system were recounted. Participants felt the advice was beneficial regardless of whether it resulted in additional income. The majority receiving additional benefits described a positive impact on life choices and control, quality of life, independence and health. Awareness of eligibility and availability of benefits and benefit advice remains low, and many older people and carers still feared the intrusion of benefit provision and advice.</td>
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<td>Dobbie and Gillespie (2009)</td>
<td>Evaluation of financial inclusion development project to improve mainstream practice, not develop new services</td>
<td>Methods included: analysis of data, meetings with project staff, 5 group interviews with service users and other stakeholders, 4 interviews with health service professionals</td>
<td>Service users valued a supported referral system to access advice and awareness of mental health issues improved amongst advice services in Glasgow over the period. Service users and carers described the impact of reduced stress and anxiety on their health and well-being. Report identifies admission to and discharge from hospital as key times at which people need help to address money problems, but they may lose existing links with service providers that operate within defined geographical areas.</td>
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<td><strong>Gillespie et al (2009)</strong>&lt;sup&gt;35&lt;/sup&gt;</td>
<td><strong>Survey of CAB debt clients</strong></td>
<td><strong>Methods:</strong> 273 questionnaires completed by CAB advisers, 10 qualitative purposive interviews</td>
<td>Changes leading to a drop in income and persistent low income were important features. Interviewees attributed stress and depression to their financial situation, regardless of the reason for being in debt. Pressure from creditors contributed to the impact debt had. Advice relieved the stress and anxiety of debt and gave some interviewees confidence.</td>
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<td><strong>Jones (2009)</strong>&lt;sup&gt;26&lt;/sup&gt;</td>
<td><strong>Longitudinal study with CAB clients in Wales to measure the long-term impact of advice-giving on clients who had approached Citizens Advice with one or more problems. Included evaluation of the health impacts of advice</strong></td>
<td><strong>Data collection began April 2007, further interviews at 6 months and at 12 months after contact with a CAB. Questionnaires incorporated SF-36 health questionnaire and Hospital Anxiety and Depression Scale (HADS) and a range of social support, demographic, and financial questions, and analysis of data on nature of advice problem(s)</strong></td>
<td>Improvements to financial wellbeing meant clients were better able to afford a range of essential items and social activities for themselves, and any dependent children. As a result of advice, clients described improvements in health: they felt better; were less anxious, less stressed and less worried about money. Some noted improved emotional states of mind and feelings of self belief. Clients with dependent children reported improved relationships and were more relaxed with their children. Statistically significant improvements were found in the SF-36 domains of Social Functioning, Role Emotional and Mental Health, and in the Anxiety and Depression scales of the HADS at Wave 2. At Wave 3 statistically significant improvements were found in the domains of Vitality, Social Functioning and Mental Health in the SF-36, and in the Anxiety scale of the HADS. Argues there is evidence of the improvement over time in clients’ health as a direct outcome of advice and support. Argues funding CAB service remains key to saving costs on health and mental health as well as improving the lives of vulnerable clients.</td>
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* These studies were included in the review by Adams et al in 2006
The Impact of Advice for Health Services and Staff

It is evidence of the associations between social problems and morbidity which supports a broad approach to service provision in general practice. Although the model of welfare rights advice based in primary care settings is now well established, provision remains patchy. An impact of this is that doctors and other health professionals are involved in the provision of advice about problems involving rights. Exploring the social problems presented to GPs, Popay et al found that the types of problems raised were most commonly concerned with welfare benefits and housing, but GPs were most likely to refer to counselling services and to a lesser extent to generic advice services. In this study doctors reported finding it difficult to keep up to date with the availability of specialist support because of the transitory nature of some services. Another study questioned the level of skills, awareness and training of health professionals to provide rights advice and advocated better measures to provide outreach advice, particularly on welfare rights, in order to promote both justice and health outcomes.

Research has identified that the stress and anxiety associated with debt can affect health and can result in visits to a GP. Drawing on this evidence, the London Health Forum highlighted that the resulting costs for the NHS could be significant, for example an hour of GP time costs approximately £160 in the city. The Forum has initiated a project to help ameliorate the health burden arising from the recession through early debt advice and other preventative measures.

The impact of partnerships with advice agencies on the host services and staff has been assessed in several studies. Greasley and Small argued that advice located in primary care was particularly valuable for practices working in areas of high deprivation where a large number of consultations have an underlying psycho-socio-economic basis. Advice work saved time and relieved the pressure on GPs and other primary health care staff in this setting. Other qualitative studies have identified outcomes including improved patient care at the same time as decreasing GP workload; a trend towards a reduction of GP consultations and new drugs prescribed, particularly amongst people whose income increased; and evidence of reduced drug-dependency. Also health care staff have reported that advice can provide a referral pathway for patients who have social welfare issues allowing them to ‘spend time more productively with patients’. Approaches to design and implementation of financial inclusion initiative are discussed in more detail below in the section on Methods of Intervention.

Approaches to researching the health impacts of financial inclusion

Within the studies identified, the lack of clear results in terms of measurable impacts on health has been attributed, at least in part, to the methods used in some studies. Issues raised include sample size, attrition rates and the relatively short period of longitudinal studies (12 months at the longest from initial to final survey or interview). This section provides a summary of issues identified concerning the research methods used. Most studies included in this review contained some discussion about methods or research design, particularly in relation to three key areas: the relevance of standard instruments such as the SF36 and HADS to assess the impact
of financial inclusion activities; the merits and demerits of control trials; and the role of qualitative and mixed-methods approaches.

**Instruments to measure the impacts of financial inclusion**

The only standard form that was used on more than two studies was the SF36 that was used in 5 studies using quantitative methods. The instruments identified in these studies are widely used to measure (self reported) health outcomes but, whilst these are well-established instruments for measuring health outcomes, they may not be appropriate for financial inclusion work. For example, Adams et al highlighted that general measures of health such as the SF36 “may not be able to pick up subtle changes in psychological and social aspects of health”.18

Greasley used SF 36 and HADS and argued that, in the absence of a randomised control trial or legitimate comparison group, the health outcomes on these standardised questionnaires, coupled with patients’ own accounts is perhaps the strongest evidence available to show that welfare advice can improve patients’ health and quality of life.34 However McIntosh et al identified a concern that some of the questions on the standard scales they used to assess health related outcomes: “may have been inappropriate for a study amongst older people and were not designed specifically for an older population”.39 They argued there is a need to look at alternative or additional measures relevant to an older population, such as maintenance of independent living. This lack of an existing ideal standard measure to assess the health, wellbeing and quality of life impacts of interventions may help to explain why so many different forms have been used in a small number of studies. It also suggests that more work is needed to develop measurements that are more suited to assessing financial inclusion interventions, addressing the situations of people who may be elderly or in poor health and reflecting a broader social agenda.

**The relevance of (randomised) control trials**

Few studies in this review involved the use of comparator or control groups and only two studies identified involved randomised control trials (RCTs), one of which was not completed because sample attrition was considered too great.39,27 A key conclusion drawn by Adams et al was that RCTs should be used in future studies to gain a clearer understanding of the health impacts of advice.18 There is a school of thought that advocates RCTs as the ‘gold standard’ in studies to assess the health impacts of an intervention. This is argued to be because they are regarded as the most reliable form of ‘scientific evidence’ since they eliminate spurious causality and bias. However, there are opposing views about the appropriateness of using RCTs as a method of assessing the impact of welfare rights advice on health, particularly where it could involve withholding information about legal rights and entitlements to benefits or income increases for the purposes of research.

Even setting aside the fact that lack of use of mainstream services is often the main reason for outreach based in health settings, others question the ethics and relevance of randomised studies and the use of control groups. Two of the co-authors in the Adams et al review were involved in a project to assess the feasibility and acceptability of a RCT of welfare rights advice and to identify appropriate health and social outcome measures in order to plan a definitive trial. MacIntosh et al
sought to minimise the effect of loss of income by having a six month delay prior to the control group being assessed for benefit entitlement. They considered this would be a reasonable compromise “between impeding the receipt of benefit entitlements and allowing time for any health improvements to appear”. However, the optimum time at which to measure any health benefits was unclear at the trial outset. They concluded that health improvements resulting from increased benefits may not be detectable after such a short time and identified that individuals in the control group did lose benefits as a result of this approach. While there is a methodological argument for delaying the welfare rights advice longer in the control group in a future trial, they recognised it is “ethically contentious, as it is hard to justify withholding an intervention that is known to be beneficial in financial terms”. 39

Abbott et al were criticised because they did not use a control trial. However, they identified the objections to RCTs as an approach for this type of study topic as including concerns about legal entitlements to welfare benefits and the fact that welfare benefits advice, in common with other financial inclusion interventions, is unlike medical treatments which are more commonly assessed using RCTs; and the resulting changes from advice or an increase in income have a value irrespective of any health effects, including the relief of poverty. For them it was:

“arguably unethical to deprive anyone, even temporarily, of money to which they are legally entitled, for the sake of an RCT when the presumed benefits to health are neither established (hence this study), nor the main outcome of the intervention. Observational studies have a valuable role to play when RCTs are not feasible.” 28

Consistent with this view, Campbell et al considered that the use of a control group was not feasible for ethical, social and practical reasons. Although they acknowledged the need for cautious interpretation of findings, they highlighted that a qualitative study was conducted alongside the quantitative research and that supported their interpretation of health and non-health impacts to some extent.24

It is difficult to envisage conditions in which the ethical concerns raised here would not arise in research to assess the impact of advice for health, wellbeing and quality of life. Advice and financial inclusion interventions are first and foremost aimed at increasing income or sometimes reducing the cost paid by people on low incomes. While there is scope to assess the effects of such interventions on health, the need to reflect the social nature of the intervention should be prioritised:

“Social interventions differ from clinical and most complex public health interventions in that changes in health are often an indirect effect rather than a primary aim of the intervention. Investigation of indirect health effects often requires choices to be made between competing values, usually health and social justice, creating a moral problem. When, as in our study, the tangible social and economic gains generated by the social interventions outweigh the theoretical possibility of marginal health effects, the moral issues are clear.” 47

Following Springett we would argue that appropriateness is intimately tied up with the nature of the object that is being evaluated.48 The question of what is being
evaluated has implications for the choice of what constitutes an appropriate framework for research and evaluation.

**Qualitative and mixed methods studies**

Mixed methods were the most common approach used in studies and those reporting concurrent qualitative and quantitative elements to studies highlighted the important contribution of qualitative research to extending understanding in what is a relatively new area of research for which the established measurement of health outcomes are less than ideal. For example, Moffatt et al highlighted how their qualitative study provided valuable information about outcome measures and helped with the overall interpretation of the concurrent pilot RCT. It enabled:

“participants to give an account of the various ways in which the intervention impacted on their lives, such as increased independence and improved quality of life, which were not explicitly measured in the pilot RCT and are challenging to capture quantitatively... The study also suggests that inclusion of a qualitative component will help to illuminate the process and outcome of a future trial.”

Similarly Winder et al described the validity and reliability of their social services based study as being “supported and strengthened through feedback within and between concurrent studies” and that primary and secondary coders (with different professional backgrounds) came to a consensus on how themes were categorised and interpreted. This approach is supported by Rogers et al who, in discussing the impact of urban regeneration on mental health, argue that:

“a combination of multi-level mode indicators and narrative accounts of individuals about mental health in the context of locality and personal changes are central for developing theories and methods appropriate for exploring the action and interaction of effects operating between structural and individual/agency levels.... Surveys which use subjective indicators of mental health and quality of life, together with qualitative research, which incorporates the narrative accounts of individuals about both the locality and personal circumstances implicated in mental health status, has the potential to assess and elaborate upon the way in which individuals perceive the effects of change."

Considered in this context, financial inclusion activity is a complex intervention and, whilst subject to more research than other areas, money advice is one aspect of potential financial inclusion activity whose effects do not necessarily follow the assumptions informing conventional assessment of health impacts. In the studies reviewed, there was some lack of satisfaction with standard measures to meet the needs of research in this relatively new field and debate around the relevance and ethics of randomisation and control trials. There were multi-disciplinary research teams in only a few of the projects identified, but where qualitative research was included in the methods used, the complexities have come through more clearly. At this early stage in a new field of research, there remains considerable scope to develop research projects from design through to analysis that reflect the complexities and multi-disciplinary nature of the issues involved.
Section 3: Models of intervention

This section draws out good practice lessons from the health-based literature to inform the development of future financial inclusion services.

Understanding the barriers or specific needs of different groups is central to effective service design. When designing financial inclusion initiatives, “what helps one group of clients may not help another”. Research has established that advice is often needed at key transition points in people’s lives such as the move into employment or job loss; when a relationship breaks down; or at the different stages in patient pathways, from diagnosis through to recovery. In addition, services need to consider how to design approaches that take account of the way that structural inequalities impact on specific groups.

Targeting groups

Income is a key predictor of financial exclusion - vulnerability to financial exclusion is related to both long-term and temporary low incomes, and to facing major unexpected expenses. Different vulnerabilities can create barriers to financial inclusion which interact with low income. What follows is a summary of those studies which have targeted an intervention to meet the needs of specific groups:

- **Older people**: It has been recognised since the early 1990s, that providing benefits advice services for older people in a health-care setting, usually a primary health-care centre, can be highly effective and beneficial. Reasons identified within the literature have included: many older people have regular contact with primary care and clinics and health centres can be less stigmatising than benefits offices. Key features of income maximisation services for older people have involved: the active creation of contacts with older people to alert them to their entitlements; pro-active offers of advice to people who are retired and particularly those in poor health, with a disability or in a caring role; and active assistance with the process of claiming.

- **Ethnic minority groups**: The review identified one project which successfully reached this target group while others have reflected on the barriers to access for this group including lack of bi-lingual staff and promotional materials. The Health Plus Project employed 6 advice workers in General Practices throughout the Bradford City Primary Care Trust (PCT) area. Most of the advice workers were from ethnic minority origins and able to provide a bi-lingual service, whilst two advice workers could access interpreters. Income generation through benefit claims was only one aspect of this project and other forms of advice and assistance (e.g. disability aids & adaptations, home care, housing, immigration) were equally important for patients. The majority of patients referred to the service were female (61%) and most were of south Asian origin. The authors suggested that the high proportion of Asian service users may reflect a greater need for help with understanding and completing welfare benefit letters and forms.
• Gender: Although the studies included in this review often reported on the numbers of men and women accessing projects there was no gendered analysis of the impact of services or health outcomes. There is increasing recognition among health policy-makers and service planners that gender is a central determinant of health status. A number of studies have explored the way men and women tend to use health and other public services differently. However, in the area of health and over-indebtedness, gendered analysis is lacking and there is potential for future research in this area.

• Families and Children: Families with young children are particularly vulnerable to poverty. However, no studies were identified that had families with children as their main focus. The recent announcement of a ‘Healthier Wealthier Children Families Financial Inclusion Project’ across NHSGGC is a welcome development in this regard. This 15 month project will target pregnant women and families with young children who are at risk of poverty. It will aim to develop expertise to address child poverty through a partnership model providing income maximisation advice and developing a strategic approach to linking the provision with health and other services.

Interventions targeted at patient groups

Health-based financial inclusion initiatives should be flexible, taking account of the needs of patients with acute conditions and provided at different stages in the patient pathway of longer term conditions. Acute, short-term episodes of ill health have a similar relationship to financial exclusion as many other dramatic shocks in people’s lives and poorer people are less likely to have a cushion to manage this kind of crisis. Here, early access to advice on benefits and action to prevent debts getting out of control during the period of illness are priorities. Different support may be needed for people with chronic health conditions: for example, where they limit ability to work, people can be denied financial goods and services as they move onto long-term low incomes.

• Cancer patients, their families and carers: In parts of the UK including some areas in Scotland, Macmillan Cancer Support has established partnerships with local money advice services to improve access to specialist advice and address the stress and anxiety associated with money worries which often arise following diagnosis for example, due to a drop in income or increased costs associated with managing their cancer. The service is delivered in places that are convenient to the clients and their families including in their own home, in hospitals and other health settings.

  o In England and Wales Macmillan Cancer Support have funded CABs to provide specialist advice to people affected by cancer. Between January 2007 and June 2009, 51 bureaux provided a Macmillan partnership service. They helped nearly 15,000 clients. The majority (54%) accessed the service via either an outreach service, home visit or at the hospital.

  o Within Glasgow, Macmillan Benefits Service was established in 2008. This is a four year partnership between Macmillan Cancer Support,
Glasgow City Council and NHS Greater Glasgow & Clyde. The service has a Glasgow-wide remit to accept referrals from anyone with a cancer-specific diagnosis and their families or carers in the local authority area. It offers a home visiting service and advisers also visit customers in hospitals and hospices if necessary. The project accepts referrals from a wide range of sources including: hospitals, hospices, health centres, and social work, family and friends and self referrals.

- **People with mental health problems and their carers:** People with mental health problems are vulnerable to financial exclusion and problem debt. In Britain they are nearly three times more likely to report debt compared with individuals without similar conditions. The Manchester Mental Health Welfare Rights Service and the GAMH financial inclusion development project are two good examples of services designed to meet the needs of this group. Evaluation of these projects has found that what works is:
  
  o Advice sessions delivered from places people with severe mental health problems go to and are comfortable in, such as day treatment centres, where user groups have drop-ins, or via home-visits if required.
  
  o Staff who are expert in benefits and advice and knowledgeable about a range of medications and mental health conditions.
  
  o Some service users require the support of their key worker or CPN to attend appointments. GAMH operates a supported referral service which puts the person referred to a money advice agency at the centre of the process. It ensures the person referred is provided with support from a worker who is experienced in supporting people with mental health problems before, during and after the referral to the money advice agency. The service also provides mental health first aid training to money advisers.
  
  o It can take several appointments for an adviser to make a needs assessment or fill forms with some clients. This may be because it takes longer for the client to build trust with the adviser, necessary to apply for key benefits such as DLA or because they have to go into hospital.
  
  o Continuity in the service provision is crucial for this client group because some can find it difficult or distressing to access advice and tell their stories to strangers.
  
  o Continuity of support through the process of admission to and discharge from hospital.

- **People affected by stroke:** A financial inclusion service was established in November 2008 by NHSGGC to meet the financial needs of people affected by stroke. Key lessons from the evaluation of this service include the need
for advice at each stage of the stroke pathway including early intervention; continuity of adviser on discharge; and holistic support covering all aspects of money advice. Unfortunately the service evaluation only focused on the money advice and income maximisation activity and lessons were not drawn on the impact or uptake of financial capability. Despite these problems with the evaluation of this, the project has a patient-centred model which aims to provide a flexible package of support and the approach to consultation and design of this service can inform the development of provision for other groups.

**Screening or referral?**

The research indicates that, whilst some projects are based on multiple referral routes (e.g. self-referral, GPs, nurses, social worker or carers), others have managed projects through tighter referral or screening arrangements to target specific groups or situations. A clear lesson from projects which have focused on access to advice through GP surgeries has been that this is an effective way for services to reach groups who are less likely to use mainstream services. However, some studies have cautioned that the primary beneficiaries of this approach tend to be older people, more than other groups at risk of exclusion and considered hard to reach, including younger families. Also the level of uptake is dependent on the attitude of practice staff towards the service, their knowledge of the service, and general promotion of the service. Popay (2007) argued that the reason that at least some GPs do not respond to patients' social problems are likely to be complex, but research suggests that they include patients' reluctance to disclose social problems and GPs reluctance to probe for these.

The most successful services tend to be where other health workers are fully supportive of the initiatives and the welfare rights advisors are an integral part of the health unit (e.g. GP practice), with other health care staff aware of appropriate clients to refer on to them. Where advice workers are more marginalised, due to location (lack of space) and/or lack of interest from healthcare staff the literature suggests the impact was less.

Hoskins et al (2005) evaluated a 15 month project where community nurses used pre-screening forms in the course of their normal daily work to screen their older clients' Attendance Allowance status. The authors argue that the pre-screening intervention overcomes the barrier of some clients' reluctance to disclose personal information and share health concerns, due to the therapeutic nature of the nurse/client relationship. Toeg et al have highlighted the challenges of this approach arguing that screening requires specialist benefit knowledge, as well as an awareness of statutory and voluntary services. He noted the educational potential that can stem from such a project in relation to all members of the primary health care team, as well as medical students and general practice registrars.

**Managing referrals and expectations**

Whilst it is vitally important that people in need of support are assisted in identifying sources of accredited advice, it is inherent in this that once directed they are actually able to access that advice: i.e. receive a consultation within a timely period. If the
number of referrals increases as a direct result of the establishment of a referral mechanism, it is imperative that the resources available to service this increase in demand are identified and released.\textsuperscript{27}

**Developing Services**

**Training needs**

Advice work in primary care and other health settings, compared to generic advice work, may require relatively longer sessions and longer-term involvement with patients to address complex problems and deal with follow-up work. To support service users with a range of complex health needs, one evaluation identified that money advisers established links with mental health organisations and accessed ‘mental health first aid’ training\textsuperscript{10}. Fitch has recommended that health staff should receive training on how to talk with patients about debt; knowing how to refer to, and support, debt advisers; but without being expected to become ‘debt experts’ themselves.\textsuperscript{5} The Royal College of Psychiatrists, with the input of a range of other organisations, has developed a guide to help health and social care workers support people with debt and mental health problems.\textsuperscript{65}

It is important to provide training for health staff about appropriate referrals. Referressees need to be aware of the range of assistance provided in order to avoid inappropriate referrals and ensure use of the service.\textsuperscript{19} A recurrent issue identified concerns health staff attitudes presenting a barrier to advice. In one study health professionals characterised claimants as ‘scroungers’ and a nurse assumed that a patient did not need DLA because he/she belonged to an ethnic minority that “looks after their own”,\textsuperscript{21} pointing to the need for equalities and diversity training to tackle barriers to referrals.

**Advertising and promotion**

To encourage engagement in any initiative, primary care staff and other groups of referrers in a number of projects have highlighted the need for feedback about referrals.\textsuperscript{15,10} Whilst it was appreciated that advice issues are confidential, feedback about the number of patients seen and general outcomes is important.

Across the literature the need for advertising and promotion of services was highlighted as an important issue. Advertisements and promotion should be visible in ‘outreach’ locations where people might be seeking help, using a service or just looking for information (e.g. Jobcentre, childcare services, libraries, GP surgeries).\textsuperscript{27} In one study surgeries with printed information available rated the adequacy of advice services more highly.\textsuperscript{65}

**Partnership working**

Good practice in delivering financial inclusion interventions in health-settings has been achieved through partnership working with local specialist advice agencies. Although research has identified that complex partnerships can limit the effectiveness of short-term projects,\textsuperscript{10} Partnerships with specialist money advice services, as well as services that are specialist in supporting groups, have been identified as effective ways to develop advice and outreach sessions and other activities that improve access to advice. For example, the
Glasgow Association for Mental Health has implemented a strategy to tackle financial exclusion and improve access to advice for people with mental health problems that involves partnership working with money advice agencies across the city, service users and carers and it is currently developing stronger links with NHSGGC.
Section 4: Conclusions and Recommendations

Conclusions

Addressing the health benefits of financial inclusion remains a complex area of intervention and research. As a very new discipline, financial inclusion and the approaches involved within it are constantly changing and developing and there remains considerable scope for further project development and research. Although this review has had financial inclusion as its focus, income maximisation work remains the central concern of most of the research identified. However, financial inclusion is not only about access to benefits and income maximisation but also debt advice, financial awareness and financial capability (how people use financial services, how they budget and pay for their bills and how they plan for the future). A growing number of health-based initiatives aiming to tackle the wider issue of financial exclusion are operating within NHSGGC and nationally, and the picture is moving fast.

Whether or not projects have involved some targeting, for example towards older people, there is limited evidence of analysis of the different situations of different groups or the impacts for groups within the target population, for example on the basis of gender, ethnic origin, disability or learning difficulties. Strategies that work for one group or situation can inform work with other groups but may not always be effective. Research and evaluation need to go beyond recording the characteristics of service users and explore different needs, impacts and outcomes of advice. However, this work should also be informed by a growing body of research on effective practice in financial inclusion work.

Investment in financial inclusion is relatively recent within the NHS, so many initiatives remain at the early stages or are short-term, and few have been evaluated for their impact on health. This presents opportunities for future evaluation and research to explore the health impacts of these approaches. There are gaps in the groups that have been targeted in initiatives and limited analysis of outcomes or analysis other than for whole populations, for example on the basis of gender. The research shows that services are effective at reaching older people, for whom primary care is a trusted source of support. While this is likely to remain important, there is less evidence of services targeting other groups at risk of exclusion, including families with children. However, a new NHSGGC project targeting this group will be well placed to contribute to building understanding of the benefits of income maximisation work in a health setting for a key group that is vulnerable to poverty.

The mental health benefits of financial inclusion activity are important in their own right and the area in which qualitative and quantitative research has shown some clear benefits. However, the wider effects of addressing the stress and anxiety of debt and low income should be explored. For example, the Mental Health Foundation has highlighted that people with anxiety disorders are at greater risk of developing a number of different physical health conditions, including coronary heart disease\textsuperscript{67}. Regardless of the causal pathways between debt and mental ill health,
the link is evident and important, particularly in view of the growing problem of personal over-indebtedness in recent years.

The review has also identified opportunities to further develop existing approaches to tackling financial exclusion. For example, in addition to welfare rights and debt advice, other linked areas of policy and practice have the potential to be considered in integrated approaches to financial inclusion because they are strongly linked to the issues of poverty and ill-health. Addressing fuel poverty is one area that has been highlighted. Prevention of homelessness and eviction and re-housing of homeless individuals is another area in which the right advice and support is essential to addressing the situation of people who are likely to have health and/ or addiction issues.

These wider issues also serve to highlight the need for a broader agenda in research that takes more account of the complexities of people’s lives. The review raised questions about whether enough account has been taken of the effects for different groups of people and different health circumstances (for example acute and chronic health conditions, mental health problems). There is some agreement that, in order to gain a better understanding of the impact on health, more sensitive measures need to be developed and studies need to be longer term than the one year that most studies have lasted in the past.

Adams et al concluded from their review that there is little need to conduct additional work to determine whether welfare rights advice has a financial effect but there was “little evidence either that welfare rights advice in healthcare settings does or does not have health and social effects, and this remains an intervention with theoretical potential to improve health”.18 We would re-emphasise the issues identified with the lack of sensitivity of standard self-report health outcome tools in the field of advice and financial inclusion impacts on health. More importantly, however, there is a need to illuminate further the evidence from quantitative research and from a body of qualitative research which supports the positive effects of benefits advice on mental health and socio-economic wellbeing. The significance of this evidence has been underplayed.

Evidence of effective practice exists, but this would benefit from further development. In particular, advice needs, like health needs, are often not static and some flexibility in service design may be needed to respond to changing needs. For example, someone diagnosed with a condition involving long-term management or a long period of recovery may have particular and different advice needs at the point of diagnosis, when entering or leaving hospital as an in-patient and during periods of recovery or deteriorating health. Such an approach would be consistent with the aim of holistic provision and the aims of providing seamless services and the use of the pathways approach. Training and information sharing are necessary components for ensuring that health and financial inclusion professionals have the right levels of awareness and expertise for the work they do.

Financial inclusion is an area of mutual concern for local government and health services and has much potential to contribute to better understanding of how services can help to reduce health inequalities. This potential also extends to addressing any unintended consequences of the way services currently work.
Recommendations: Research

**Recommendation 1:** More research is needed to broaden understanding of the extent to which financial exclusion and inclusion are affected by factors such as gender, family circumstances, age, ethnicity or disability and their relationship to the impacts of financial inclusion initiatives for health, wellbeing and quality of life for different groups within target populations.

**Recommendation 2:** Financial inclusion is a complex intervention. There is a great deal of good work being done that is not necessarily providing the depth of research and evaluation that is still required. There is considerable potential for financial inclusion initiatives to contribute to an agenda for improving health. However, for its contribution to be understood better and the social impacts of financial inclusion taken into account more fully, there is a need for:

- Multi-disciplinary research involving people with expertise in both health and financial inclusion
- Mixed-method approaches to research that draw upon the experiences of individuals to develop more effective methods for assessing the impact of financial inclusion interventions in relation to health, wellbeing and quality of life. Such work should build upon, but not rely on, existing standard health outcome measures such as the Short Forms that have been identified as insufficient for financial inclusion initiatives and for research with some groups of people, including older people
- Longitudinal research is needed, particularly for improving knowledge about the impacts of financial inclusion interventions for physical health and condition management. Ideally this needs to go well beyond the one year research period of most existing studies

**Recommendation 3:** Further research is needed on the links between debt and mental health, particularly to increase understanding of the ways in which financial inclusion activities can alleviate mental health issues and the potential for such interventions to contribute to reducing the physical health risks associated with poor mental health.

Recommendations: Practice

**Recommendation 4:** Monitoring of the characteristics of people using financial inclusion services should be kept under review to ensure that it can reflect both project and wider policy priorities. For example, monitoring should include family situation / relationship status, having dependent children and caring roles in addition to the social characteristics that tend to be recorded more routinely.

**Recommendation 5:** Evaluation and development of financial inclusion projects and services should take account of the effectiveness of services at reaching different groups, including within target populations. For example, do services targeting older people effectively meet the needs of both women and men?
**Recommendation 6:** Existing research and evaluation reports provide practical guidance on improving access to advice for people in a wide range of situations and this body of work can do much to inform development and review of projects and services, including adapting existing effective practice to improve service access for other vulnerable or disadvantaged groups.

**Recommendation 7:** Projects and services should be developed in a way that recognises the importance of responding to changing needs over time. This is consistent with holistic service provision and the pathway of care approach.

**Recommendation 8:** In view of the importance of a warm home, particularly for young and older people, NHSGGC should consider how addressing fuel poverty can be incorporated within its approach to financial inclusion. Linked to this, housing circumstances, particularly the risk of or actual homelessness, are potentially important areas for advice and support, particularly given that people in this situation are likely to have health or addiction issues.

**Recommendation 9:** The wider issues of fuel poverty and homelessness highlight that there may be a need for wider links, for example with services addressing advice on homelessness and benefits, including CABs and Shelter, in a broad agenda to tackle financial exclusion.

**Recommendation 10:** Partnership working needs to include key health and financial service providers, but also service users and carers and the services that support them can provide valuable inputs to project development (for example, key workers). This could include development of alternative approaches to delivering financial inclusion advice, such as peer advice and support. In some circumstances, the input of other service providers may be important, such as housing providers and domestic fuel suppliers.

**Recommendation 11:** Training, awareness raising and capacity building should be kept under review to take forward financial inclusion work. This is likely to involve different training need for different groups and services, for example:

- For mainstream financial inclusion services training may need an emphasis on health needs and their impact on access to services and financial inclusion, with additional training where necessary, such as mental health first aid for money advisers
- For health service staff training should raise awareness of general rights and entitlements and the specific or general sources of help, including encouragement towards referral rather than seeking to gain detailed knowledge of benefits or other financial inclusion interventions
- For all groups, equality and diversity training may be important, particularly in projects involving screening of potential clients for financial inclusion interventions
NHSGGC Financial Inclusion Group will consider which recommendations can be taken forward through its review of its action plan for 2010/11 and which are longer term. There are opportunities for many of the recommendations to inform planning and service provision (e.g. Children & Families Project, Mental Health services, Long Term Conditions, Older People); development of guidance (e.g standards group; training initiatives) and monitoring (e.g. health benefits research group; policy framework indicators).
Appendix 1: Search Strategy and Analysis

To ensure that the relevant evidence was gathered within the timescale, we undertook an approach that is in accordance with some of the best practices involved in Rapid Evidence Assessment (REA). This approach ensures that rigorous and transparent search and analysis procedures produce a valid, robust and reliable analysis of the field. It also draws out lessons for future research and practice.

The key principles and steps in the REA method which can be applied in this project include the following:

- Developing an informed and systematic search strategy using electronic and print sources to ensure comprehensiveness within the constraints of the timetable
- Establishing explicit inclusion criteria for identifying relevant sources
- Assessing the relevance of identified articles in accordance with a standardised protocol
- Critically appraising evidence gathered to sift out studies of poor quality in accordance with explicit and transparent criteria
- Collating descriptive and analytical outlines of the selected evidence in a systematic and standardised format
- Providing an accessible and coherent narrative overview of the evidence and conclusions which follows logically from the analysis process

This appendix reports on the key elements in relation to the three key stages of this review which include the following:

- Stage 1: Data Search
- Stage 2: Data Analysis

Stage 1 – Data Search

The first stage initially involved clarifying the parameters and relevant search criteria to guide the literature review and evidence gathering processes. The agreed set and range of search terms is presented in Table A1 below. These keywords were identified based on understanding of the theory, research and debates in the area, and specified in advance to ensure that the search process and criteria for including and selecting studies is explicit and defendable.

The following points provide an overview of the data search:

- **Searches of electronic databases**: the main source of information on social policy and related research are bibliographic databases - collections of document summaries, usually with abstracts and some subject indexing based on a consistent coding procedure. The nature of coding and classification of content is often specific to each database and so the keywords in table A1 were used to search the thesaurus of each of the following databases:
<table>
<thead>
<tr>
<th>AND</th>
<th>AND</th>
<th>AND</th>
</tr>
</thead>
<tbody>
<tr>
<td>Banking</td>
<td>Inclusion</td>
<td>Health</td>
</tr>
<tr>
<td>Credit</td>
<td>Exclusion</td>
<td>Mental health</td>
</tr>
<tr>
<td>Debt</td>
<td>Poverty</td>
<td>Primary care</td>
</tr>
<tr>
<td>Money</td>
<td>GP (General Practitioners)</td>
<td>Research</td>
</tr>
<tr>
<td>Money advice</td>
<td>Social work</td>
<td>Outcome</td>
</tr>
<tr>
<td>Over-indebtedness</td>
<td>Hospitals</td>
<td>Impact</td>
</tr>
<tr>
<td>Saving (s)</td>
<td>Nurse</td>
<td>Result</td>
</tr>
<tr>
<td>Income maximisation</td>
<td>Effect</td>
<td></td>
</tr>
</tbody>
</table>
| Welfare benefits | Satisfactio
| Insurance   | Cost        |
| Debt advice | Benefit     |
| Affordable credit | Review |
| Loans       |             |
| Financial services |             |
| Credit Union |             |
| CDFI        |             |
| Unbanked    |             |
| Financial   |             |
| Bank account |             |

- ABI/ Inform
- Applied Social Science Index and Abstracts (ASSIA)
- Cambridge Scientific Abstracts: Worldwide Political Science Abstracts
- Cumulated Index to Nursing and Allied Health Literature (CINAHL)
- Education Resources Information Centre (ERIC)
- EconLit
- International Bibliography of the Social Sciences
- Planex
The search strings used to search each database along with a record of the number of ‘hits’ are included in this document in table A2 below.

A. **Hand searches of specific journals:** the electronic contents pages of Health and Social Care in the Community (Volumes 6–12, 1999–2009), BMC Public Health (Volumes 1-9, 2001-2009) and the Journal of Social Policy (volumes 28–38, 1999–2009) were scanned to identify relevant publications. These journals were chosen because of their relevance to the subject area and the perception that substantial relevant work had been published in them.

B. **Searches of internet search engine:** searches were made of the internet search engine Google [http://www.google.com](http://www.google.com) and Google Scholar [http://scholar.google.co.uk/](http://scholar.google.co.uk/) using the keywords in Table A2. The first 100 results returned by each search strategy were scanned for relevance and those judged to be potentially relevant followed up.

C. **Suggestions from experts and those working in the field:** 'Experts' – identified as such either by frequent publication in the area, or through personal contacts of the research team – were contacted directly and asked for help with identifying relevant literature or providing further contacts. Relevant reports were picked up from e-bulletins from a range of relevant organisations including Social Policy Digest, Transact and The Information Hub.

D. **Local monitoring and evaluation reports and reviews:** In addition to the review, the researchers have undertaken analysis of local monitoring and evaluation reports and reviews conducted by other health services across the UK. The commissioners have provided the research team with a number of relevant documents for this stage of the process.

### Table A2: Database Searches

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<td>CINAHL</td>
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<td>2</td>
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<td>PLANEX</td>
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<td>37</td>
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<tr>
<td>ERIC</td>
<td>Mental-Health.DE. OR Mental-Health.DE. OR Health-Needs.DE. OR Health-Promotion.DE. OR Public-Health.DE. OR Health-Personnel.DE. OR Community-Health-Services.DE. OR Mental-Health-Workers.DE.) AND (Debt-Financial OR Banking.W..DE.</td>
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<td>Relevant Articles</td>
<td>Results</td>
</tr>
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<tr>
<td>Psychinfo</td>
<td>DE=(&quot;banking&quot; or &quot;financial strain&quot; or &quot;social security&quot;) and (&quot;poverty&quot; or &quot;social equality&quot;) and (&quot;mental health&quot; or &quot;primary health care&quot; or &quot;social casework&quot; or &quot;therapeutic processes&quot; or &quot;well being&quot;)</td>
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<tr>
<td>ABI/ Inform</td>
<td>&quot;banking&quot; or &quot;debt&quot; or &quot;unbanked&quot; or &quot;credit&quot; or &quot;savings&quot; or &quot;money advice&quot; or &quot;welfare benefits advice&quot; or &quot;income maximisation&quot;) AND (&quot;health&quot; or &quot;mental health&quot; or &quot;social work&quot;) AND (&quot;inclusion&quot; or &quot;exclusion&quot; or &quot;poverty&quot; or &quot;income inequality&quot;) AND (&quot;evaluation&quot; or &quot;study&quot; or &quot;pilot&quot; or &quot;research&quot; or &quot;outcome&quot; or &quot;impact&quot; or &quot;result&quot; or &quot;effect&quot; or &quot;satisfaction&quot; or &quot;cost&quot; or &quot;benefit&quot; or &quot;review&quot;) AND PDN(&gt;1/1/1990) AND NOT AT(book review) AND NOT (fiscal)</td>
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<td>Applied Social Science Index and Abstracts (ASSIA)</td>
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<td>Cambridge Scientific Abstracts: Worldwide Political Science Abstracts</td>
<td>(DE=(&quot;debts&quot; or &quot;banking&quot; or &quot;bankruptcy&quot; or &quot;budgets&quot; or &quot;consumer law&quot; or &quot;credit&quot; or &quot;finance&quot; or &quot;financial support&quot; or &quot;income&quot; or &quot;insurance&quot; or &quot;loans&quot; or &quot;money&quot; or &quot;saving&quot; or &quot;welfare recipients&quot;)(DE=(&quot;health professions&quot; or &quot;community mental health&quot; or &quot;health care services&quot; or &quot;mental health&quot; or &quot;mental health services&quot; or &quot;mental illness&quot; or &quot;public health&quot; or &quot;social services&quot; or &quot;stress&quot;))</td>
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</table>

34
E. Searches of specific websites: the websites of a number of specific organisations that sponsor and conduct financial inclusion and social policy research are being searched on an ongoing basis and include:

- Government (UK)
- HM Treasury
- Cabinet Office
- Ministry of Justice/Department for Constitutional Affairs
- Department for Work and Pensions
- Department of Health
- Department for Business, Innovation and Skills
- Communities and Local Government
- Legal Services Commission/Legal Services Research Centre
- Scottish Executive
- Welsh Assembly
- Other bodies
  - NHS Health Scotland
  - Financial Services Consumer Panel
  - Financial Services Authority
  - Joseph Rowntree Foundation
  - Runnymede Trust
  - Institute for Fiscal Studies
  - Centre for Research on Social Policy
  - New Policy Institute
  - Friends Provident Foundation
  - Citizens Advice
  - Money Advice Trust – Information Hub
  - Sainsbury Centre for Mental Health
  - Mind
  - Macmillan Cancer Support

- Academic
  - ESRC
  - Personal Finance Research Centre
  - Research Unit for Financial Inclusion

F. Reference lists from relevant studies: the reference lists of all studies assessed to be relevant were scanned to identify other relevant work, as were the reference lists of previous reviews in this area.
Stage 2 – Data Analysis

To ensure consistency in analysis, each source identified and judged relevant in the literature search was reviewed using an analytical template covering:

- Nature of the study, article or report: e.g. original quantitative, qualitative or mixed-method research; policy commentary; theoretical analysis; research summary, etc.
- Specific subject/topic; aims of the study
- Relevance to and implications for health impact of financial inclusion activity and/ or practice in health services
- Further sources of information or reading identified

The evaluation of the quality of research studies and validity of evidence has been informed by our existing analytic expertise and familiarity with professional standards. For example, the Social Policy Association published a framework applicable to assessments of quantitative research studies, which included the following appraisal criteria (Becker et al., 2006):

- Validity: correspondence between data and conceptualisation; fitness for purpose of the research method
- Reliability: robustness of research instruments and consistency of observations
- Replicability: explicitness and transparency of method, likelihood of generating similar results
- Generaliseability: application of findings to wider cases or population

Following the appraisal of relevance and quality, the selected studies were analysed systematically in relation to content. A code list of emerging themes from studies was generated and iteratively revised in the light of new sources and results. The analysis has identified any recurring issues and findings from the literature (e.g. the frequency of particular outcomes or associations), and comment on these in terms of the relative strength of evidence and quality of the studies reporting these conclusions.
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Firth L (2009) *Funding ideas to overcome the impact of ill health on financial inclusion.* Friends Provident Foundation.


Mably S & Jones N (2009?) *The Links between Over-Indebtedness and Health in Wales*, Wales Centre for Health


Halliwell E (2009) In the face of fear: How fear and anxiety affect our health and society, and what we can do about it. London: Mental Health Foundation