The stigma of claiming benefits: a quantitative study

Abstract
Stigma has long been viewed by some as essential to discourage excessive claims, yet seen by others as a cause of non-take-up by people in need and as a form of symbolic violence. More recently, there has been a resurgence of interest in the links between shame and poverty (including the role of benefits), and particular concerns about media/political rhetoric in the UK. Yet while our knowledge of benefits stigma has been enhanced by theoretical/qualitative contributions, few quantitative studies examine its extent or patterning. This paper therefore reports the results of a 2012 nationally-representative survey in the UK. It finds sub-types of stigma are reported by 10-19% per benefit, but 34% report either personal stigma (their own view) or stigmatisation (perceived stigma by others) for at least one benefit, and over one-quarter say a stigma-related reason would make them less likely to claim. One-third of claimants themselves report some degree of stigma around their claim. Against the predictions of ‘dependency culture’ claims, however, respondents in high-claim areas were more likely to stigmatise benefits, both before and after accounting for other factors. The paper concludes by considering lessons for future benefits stigma studies, and policy options to reduce benefits stigma.

Key words
Benefits; social security; welfare; stigma; shame; take-up; survey

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Introduction

Stigma has long been central to debates around social security. Some have always seen stigma as essential in discouraging excessive claims; this was the logic of the Victorian workhouse, is implicit in the economic tradition of analysing stigma (Besley and Coate, 1992), and continues today, with the right-wing American commentator Charles Murray arguing that ‘stigma makes generosity feasible’. Others see stigma as a cause of non-take-up by the very people that benefits are designed to help, and moreover as a stark injustice in its own right. In a much-cited passage, Robert Pinker (1971:175) went as far as arguing that ‘The imposition of stigma is the commonest form of violence used in democratic societies … [It] can best be compared to those forms of psychological torture in which the victim is broken psychically and physically but left to all outward appearances unmarked.’ Versions of these debates can be found around poverty relief for centuries (Golding and Middleton, 1982; Page, 1984; Stone, 1984; Spicker, 1984).

Overlaid on this debate, though, there has been a more recent resurgence of interest. This has partly been because of an increasing focus among international poverty scholars on the ‘psychosocial’ dimensions of poverty, including disrespect, embarrassment and shame (Jo, 2013; Pemberton et al., 2013). A cornerstone has been Amartya Sen’s argument that the avoidance of shame is not just part of a relative definition of poverty, but also part of an absolute conception of poverty (Sen, 1983:167). There is evidence that the shame of poverty is globally universal (Walker et al., 2013), and that benefits stigma itself has been found to contribute to it (Walker et al., 2013:226; Jo, 2013:524; Pemberton et al., 2013:26), and we return to the relationship of shame and stigma below. Such findings have led the International Labour Organization to recommend that social protection floors have ‘respect for the rights and dignity of people covered by the social security guarantees’ (Gubrium and Lødemel, 2014:212).

A further recent resurgence of interest has come in the UK, given a perception that benefits have become increasingly stigmatised due to a media and political discourse of ‘scrounging’ (Baumberg et al., 2012). This has been argued by UK charities, several of whom argued in 2012 that ‘this narrative of benefit scroungers….does risk stigmatising all people with a disability’;² has been suggested in the academic literature (Taylor-Gooby, 2013); and claimants themselves report that this discourse makes them feel stigmatised (e.g. Garthwaite, 2014; Walker et al., 2013:226). This does not mean that benefits stigma is only of interest in 21st-century Britain; rather, it seems to be an endemic feature of most social security systems (see below), that develops into epidemics in particular circumstances.
Yet what has been missing through these debates is the quantification of benefits stigma; rarely for an important social policy issue, little quantitative evidence exists, and that which does (reviewed below) is largely dated and partial. Quantification undoubtedly requires simplifying and reshaping any object of study, putting dissimilar phenomena in the same conceptual box in order to count them; hence qualitative evidence is crucial (in the US, e.g. Seccombe et al., 1998; in the UK, e.g. Garthwaite, 2014; Shildrick and MacDonald, 2013). However, the lack of quantitative evidence prevents us from asking basic questions such as: how common really is benefits stigma in contemporary Britain? And is it true that there is a ‘dependency culture’ of low stigma in areas of high claims?

This paper reports the results of a nationally representative survey in Britain designed to answer such questions. Before this, however, the paper considers what benefits stigma is, what impacts it has, and how best to measure it.

Understanding benefits stigma

We define stigma as a situation “when a person possesses (or is believed to possess) ‘some attribute or characteristic that conveys a social identity that is devalued in a particular social context” (Crocker et al 1998, cited by Major and O’Brien, 2005:394-5) – in this case, claiming benefits. This widespread definition dates back to Erving Goffman’s influential work on managing potentially discreditable attributes (Goffman, 1963), much as some of Goffman’s views (particularly his treatment of disability stigma) have since been superseded (e.g. Major and O’Brien, 2005). Within this, we must further split ‘personal stigma’ from ‘stigmatisation’ (Chase and Walker, 2013; Taylor-Gooby, 1976:36; Stuber and Kronebusch, 2004). ‘Personal stigma’ is a person’s own feeling that claiming benefits conveys a devalued identity. In contrast, ‘stigmatisation’ is the perception that other people will devalue your identity. There is no necessity for these to overlap; claimants can think that other people look down on claiming, while personally thinking that there is nothing to be ashamed of. There is also stigma in the process of claiming benefits, from a combination of the lack of privacy involved (particularly in means tests and cohabitation tests), the demeaning experience of long waits, and feeling looked down on by suspicious staff (Finn et al., 2008:45-6). We refer to this as ‘claims stigma’ (see also Walker, 2005:197), and it often emerges in qualitative research as the most powerfully-felt stigma of all (Pemberton et al., 2013:28).

In terms of why claiming benefits might be stigmatised, the predominant view (Stuber and Schlesinger, 2006; Spicker, 1984; Pinker, 1971) is that stigma is rooted in norms of reciprocity. People who fail to reciprocate gifts incur social penalties, and to the extent that benefits are perceived to be gifts, they will be stigmatising. Beyond this, though, the degree of stigma depends upon whether claimants are perceived as ‘deserving’ recipients. There is extensive evidence that
people make deservingness judgements based on *inter alia* claimants’ level of need, whether they are seen as blameworthy, and their gratitude for the gift (van Oorschot, 2000) – and almost by definition, claimants seen as ‘undeserving’ have a devalued social identity. This helps explain why benefits stigma will vary across benefits and countries: stigma will be lower where claimants are seen as entitled to their benefits, and (where entitlement is weaker) where claimants are seen as more deserving. As Albrekt Larsen (2006) has argued, it is contributory or citizenship-based systems that tend to close down issues of deservingness (compared to means-tested systems) and which are therefore likely to have lower stigma.

This conceptual basis also allows us to understand the relationship between the closely related concepts of ‘stigma’ and ‘shame’. Shame is a powerful emotion involving a negative assessment of the self compared to one’s own expectations and the imagined expectations of others (Chase and Walker, 2013). It is likely to occur in social interactions where others (e.g. benefits officers) behave as if benefits claiming is stigmatised, and more broadly, it is a possible consequence of seeing claiming benefits as a devalued status. However, stigma may not lead to shame where people identify strongly with the identity in question and contest others’ devaluation (Major and O’Brien, 2005), or if people consider *themselves* a deserving claimant, and that the stigma only applies to undeserving others. This has been documented in both the US (Seccombe et al., 1998) and Britain, where claimants often regard themselves as deserving claimants while complaining about the ‘scrounging’ of others (Garthwaite, 2014:12; Shildrick and MacDonald, 2013:301; Dean and Taylor-Gooby, 1992; Chase and Walker, 2013). As we shall see, this has important implications for understanding the extent and consequences of stigma.

The extent of benefits stigma

While we therefore have some understanding of benefits stigma, there is only a limited literature that has attempted to quantitatively measure its scope. Where surveys of claimants of means-tested benefits have been done, around half report feeling ‘embarrassed’ (Golding and Middleton, 1982) or ‘bothered’ (Cox, 1968) about claiming. Fewer surveys focus on how claimants perceive others think, but 19% of Danish and 34% of Finnish long-term unemployed people report feeling that ‘people looked down a little’ on them because of their claim (Albrekt Larsen, 2006, Table 7.2). Recent UK evidence also suggests that around a quarter of claimants have hidden their claim at some point (Who Benefits?, 2013).

There are similarly few studies of the levels of stigma held by the general public. Those that exist find that $\frac{1}{2}$ to $\frac{2}{3}$ of their samples would feel ‘embarrassed’ about claiming (UK; Who Benefits?, 2013) or ‘do not want other people to know’ (US; Stuber and Kronebusch, 2004); that people ‘tend to look down’ on claimants (Scandinavia; Albrekt Larsen, 2006 Table 6.2) or ‘don’t respect’ them (US; Stuber
and Kronebusch, 2004); or that there is ‘a stigma’ to claiming (UK; Breese, 2011). However, only one-in-ten say they would feel sufficiently ashamed to keep their claim a secret from their friends and family (UK; Turn2us, 2012). Similarly high proportions of people express claims stigma: just under half of low-income US families agreed that ‘the application process for welfare is humiliating’ and that ‘many people are poorly treated when they apply’ (Stuber and Kronebusch, 2004), and similar proportions of British people in 2000 agreed that ‘people receiving social security are made to feel like second class citizens’. This suggests that benefits stigma is widespread, even if many claimants do not feel stigmatised or ashamed. However, there are a number of problems with these studies: they were either conducted in the 1960/70s using small, unrepresentative samples (Golding and Middleton, 1982; Williamson, 1974), are only asked to claimants rather than the general population (Handler and Hollingsworth, 1969; Cox, 1968), or are low-quality advocacy polls (Turn2us, 2012; Who Benefits?, 2013). Two recent studies (Albrekt Larsen, 2006; Stuber and Kronebusch, 2004) are particularly high-quality, but neither distinguishes ‘personal stigma’ from ‘stigmatisation’, nor compares stigma among claimants vs. non-claimants, nor examines whether there is any evidence of a ‘dependency culture’. Building on both of these studies in our own survey, we return to these issues below.

Stigma and benefits take-up

A more common way of looking at stigma is to see if people give shame-related reasons for non-take-up. Perhaps against expectations, though, these studies consistently show that shame/stigma are only cited by 5-25% of eligible non-recipients, who instead predominantly say they are not eligible or do not need benefits (e.g. Daponte et al., 1999; Taylor-Gooby, 1976). Qualitative evidence similarly suggests that stigma ‘was almost meaningless’ for people in desperate need (Rogers-Dillon, 1995:455), while polling of the general public finds relatively small numbers saying that ‘I would never claim welfare benefits, even if needed, as I couldn’t live with the shame’ (4% in Turn2us, 2012) or ‘I would feel embarrassed and therefore would not claim [benefits]’ (12% in Who Benefits?, 2013).

While some therefore believe that stigma is not a major driver of take-up (Spicker, 1984:65), another explanation is that stigma is being under-reported as ‘the admission of stigma is itself stigmatising’ (Taylor-Gooby, 1976:37), with some respondents instead using ‘colloquialisms of shame’ like ‘embarrassment’ (Chase and Walker, 2013:5), and others not admitting to negative feelings at all. Supporting this view (although see Daponte et al., 1999), some non-recipients continue not to claim after being told they are eligible (Taylor-Gooby, 1976), and some claimants’ descriptions of not feeling eligible seem closely linked to stigma (Garthwaite, 2014:9). Moreover, ‘it might be the factor of stigma which prevents potential claimants from absorbing the information which is presented to them about means-tested benefits’ (Lister, cited by Golding and Middleton, 1982:161). Even if the
role of stigma remains uncertain, it is clear that means-tested systems generally have lower take-up (van Oorschot, 2002). In our survey below, we provide updated evidence on shame as a stated reason for non-take-up.

‘Dependency culture’
Our survey enables us to investigate one further issue: the ‘dependency culture’ hypothesis. This does not just argue that claimants themselves do not stigmatise benefits, but that there are groups who collectively do not do so. For example, Charles Murray (1990/1996:26) has argued that the values of the British underclass ‘are now contaminating the life of entire neighbourhoods—which is one of the most insidious aspects of the phenomenon, for neighbours who don’t share those values cannot isolate themselves’. These ideas continue to be found within public debate; the Channel 4 television programme ‘Benefits Street’ in 2014 generated a series of headlines about the social norms in a street ‘where 90% of the residents are on benefits’. More credibly, unemployment has been found to have weaker impacts on wellbeing in high-unemployment areas (Clark, 2003), and economic evidence convincingly shows that people are more likely to claim benefits if the people around them also claim (Bertrand et al., 2000; Rege et al., 2012).

However, as these researchers admit, these studies cannot disentangle stigma from information effects. Moreover, both qualitative studies and surveys have consistently failed to find a reduced work ethic among groups who are more likely to claim benefits (e.g. Shildrick and MacDonald, 2013; Dunn, 2011; Dean and Taylor-Gooby, 1992); indeed, lower-educated people are more likely than those better-educated to say that a ‘bad job’ is preferable to unemployment (Dunn, 2011). Yet while these studies have made strong critiques of the dependency culture thesis, there is one claim that has not been directly tested: that people in high-claim areas have lower benefits stigma. The present study therefore offers a valuable opportunity to contribute direct, nationally-representative evidence to this debate.

Methods
We commissioned a bank of questions within an IpsosMORI face-to-face omnibus survey in May 2012, using quota sampling: Primary Sampling Units were randomly selected within relatively homogeneous ACORN area types, and interviewers then had to meet a quota by age, gender and working status within these streets. The resulting sample is then weighted by age, social grade, region and working status (within gender) according to a random probability sample. While not as desirable as full random sampling, it is nonetheless more likely to be nationally representative than nearly all previous surveys of benefits stigma, and also than the internet poll samples that are now becoming common in public attitude research. Given that the attitudes of benefit claimants
themselves were of particular interest, we added a ‘claimant boost’ from two further waves; the total sample was 2,601, of whom 1,123 have claimed benefits/tax credits in the past year (results are reweighted to maintain representativeness).

In designing the questions for our new survey, we reviewed previous questions, but faced the problems that (i) almost none of these questions have been validated; (ii) nearly every study uses different questions; and (iii) none of the previous surveys differentiate between personal stigma and stigmatisation. We therefore designed new questions, building on previous studies and with reference to the wider literature on measuring stigmas such as disability (Van Brakel 2006) and poverty (Reyles, 2007). Questions were both reviewed by our advisory group and underwent a small (n=4) cognitive testing.

Our first set of questions focused on stigma itself. This began by investigating personal stigma, asking ‘How much do YOU YOURSELF agree or disagree, that people should feel ashamed to claim…’, and then presenting the following five types of benefits in a random order:

- “…in-work Tax Credits? (These are wage top-ups for the low paid)
- …Jobseekers Allowance? (This is for unemployed people)
- …Employment and Support Allowance (This is for people whose sickness or disability limits their ability to work)
- …Income Support for single parents?
- …Housing Benefit? (This is help with rent for people on a low income)”

We decided to ask about being ‘ashamed’ rather than ‘embarrassed’ given recent academic debates around shame – note however that these are not questions about shame itself (whether respondents felt ashamed), but are questions about stigma (whether people should feel ashamed, as an indicator of a devalued social status). Respondents were asked to express agreement on a 0-10 scale, where 0 was labelled ‘strongly disagree’, 5 was labelled ‘neither agree nor disagree’, and 10 was labelled ‘strongly agree’. For analysis, we recoded 0-4 into ‘disagree’ and 6-10 into ‘agree’.

We then looked at stigmatisation, asking ‘How much do you think PEOPLE IN GENERAL in Britain would agree or disagree, that people should feel ashamed to claim [benefit]’, using the same list of benefits and response categories. Thirdly, to capture claims stigma, we invited people to agree/disagree (again on a 0-10 scale) with the statement ‘People are generally treated with respect when they claim benefits’. This question was deliberately worded in the reverse direction for balance, a decision we return to in the Conclusion.
Our second set of questions looked at the link between stigma and take-up, and the role of shame (people’s own feelings) within this. Non-claimants were asked, 'Which of the following, if any, would make YOU YOURSELF less likely to claim [tax credits/benefits], if you thought you needed and might be entitled to them?’ [emphasis in questionnaire]. Past-year claimants were instead asked, 'Which of the following, if any, have made YOU YOURSELF delay or not claim benefits in the past, from the point you needed and thought you might be entitled to them?’ (There is evidence that non-take-up is often about delays in claiming (Craig, 1991:543-4)). Note that respondents were instructed to ignore issues about perceived need/eligibility, given the debate about whether this is stigma-related. Each respondent was asked two versions of this question, firstly about in-work tax credits and then about out-of-work benefits.

For both versions, respondents were given the following options, which we group into four categories:

1. Category: Personal shame: • How you would feel about yourself for claiming (e.g. ‘pride’, dislike of ‘charity’) • Thinking [benefits/tax credits] are for other people, not people like me
2. Category: Social shaming: • How family, friends or neighbours would react
3. Category: Claims shaming: • Having to provide personal information (about income or having a partner) • How I would be treated by officials while applying
4. Category: Non-shame reasons: • Too hard to figure out if I’m entitled • Too much hassle to apply for them • Another reason – please specify

The survey further asked about a number of socio-demographics factors that are used as controls below: gender, age group (16-24/25-34/35-44/45-54/55-59/60-to-pension age/older), presence of children, marital status (grouped into married/civil partnership/living together, single, widowed/divorced/separated), employment status (full-time/part-time/not working), tenure (owner-occupier/social housing/other), qualifications (degree/A-level/less than A-level/none) and own benefits claims (out-of-work benefits/in-work tax credits/other benefits). Unlike previous stigma studies, we also merged-in contextual data linked to people’s home postcode. At the neighbourhood level (Lower Super-Output Areas), this included out-of-work benefit claims (as a share of the 16-64 population); at the local authority level, this included unemployment (as a share of the working-age population) and migration rates (the proportion of people born in non-western countries). Further details on the derivation and descriptive statistics of all variables are given in Web Appendix WA1. The data (and Stata code) are openly available for other researchers’ use at www.benbaumberg.com/publications.
Results

Levels of benefits stigma

Respondent’s reported level of stigma is reported in Table 1 below. Looking at the maximum stigma reported for any benefit (‘All’), 20.4% *personally* think that people should feel ashamed to claim at least one benefit (‘Personal stigma’), while significantly more (27.2%, p<0.01) believe *others* feel that people feel ashamed to claim at least one benefit (‘Stigmatisation’). A further group of almost the same size said they ‘neither agreed nor disagreed’ (14.4% for personal stigma, 20.3% for stigmatisation; see Web Appendix WA2 for the full table and responses on the ten-point scale), which may indicate a further group of people who think that claiming is stigmatised but to a lesser extent.

Claimants sometimes – but not always – perceive less stigma than the general population (due to difficulties of comparability when aggregating across different types of benefit, we focus on individual benefits). Claimants of unemployment and incapacity benefits were as likely as others to believe they should feel ashamed to claim (about 10% agreeing), but claimants of single parent benefits, housing benefit or in-work tax credits were less likely to agree they should feel ashamed (6-7% agreeing vs. 10-12% of non-claimants; p=0.01, 0.05 and 0.01 for each benefit respectively). Levels of stigmatisation, however, were similar between claimants and the general population for each benefit. Overall, a sizeable minority of benefit claimants themselves think that people should be ashamed to claim their benefit (8.7% of claimants) or think that people in general feel they should be ashamed (20.6%), which *potentially* – if they do not deflect this (see above) – may lead them to feel shame.

Previous studies have shown marked differences in perceptions of stigma for different benefits - ‘welfare’ is considerably more stigmatised than Medicaid or social security in the US (Stuber and Kronebusch, 2004; Williamson, 1974), and in Sweden, 68% thought that people often look down on the recipients of social assistance, compared to only 19% for unemployment benefit and just 4% for the basic pension (Albrekt Larsen, 2006). It is therefore surprising that there is so little difference in stigma across the five benefits here. There were statistically significant differences for both personal stigma and stigmatisation, with stigma being lowest for incapacity benefits and particularly tax credits, and greatest for single parent benefits. However, the extent of the differences in Table 1 is relatively small. We return in the Conclusion to ways in which differences between benefits could be explored further in future research.

[Table 1 about here]
When we turn to claims stigma, levels increase considerably – as many as 57.8% of people disagreed that ‘People are generally treated with respect when they claim benefits’ (with only 20.8% agreeing), which is similar among claimants and non-claimants (57.1% vs. 60.0% disagreeing). Surprisingly, the association of claims stigma to personal stigma and stigmatisation is negative. For example, those who reported personal stigma were much less likely to report claims stigma (40.8% vs. 20.8% agreed claimants are treated with respect; weighted χ² test p<0.001). This suggests that the more stigmatising respondents think benefits should be, the less respectfully they think people should be treated, and the more likely they are to say that this (lower) threshold is met.

**Stigma and benefit take-up**

The second set of questions asked non-claimants if shame would affect their take-up of benefits if they were eligible and in need, or asking claimants if shame had stopped/delayed them claiming in the past; the results are shown in Table 2. Combining the two question versions, 16.9% said that their benefits take-up would be affected by personal shame (pride, thinking benefits are for other people), 12.8% by claims shaming (personal information or treatment by officials) and only 4.3% by social shaming (how other people would react). In total, 26.6% of people gave at least one shame-related reason for not claiming.

Comparing the type of benefit claimed, we see that people are slightly less likely to say that shame influences their decision to claim tax credits than out-of-work benefits (26.6% vs. 24.1%, p=0.02) – but that this is only the case for personal shame (16.9% vs. 13.0%, p<0.01), and not social shaming or claims shaming. The results are even more striking when we focus on the past experiences of claimants themselves. Only 14.1% of tax credit claimants gave a shame-related reason for delaying or not-claiming tax credits in the past, compared to 22.8% of benefits claimants.

[Table 2 about here]

**Overlaps between stigmas**

It might be expected that respondents are divided between those reporting no stigma and those reporting multiple types – but this is only partially the case, as shown in Table 3. Even just comparing personal stigma and stigmatisation, 6.8% of respondents reported personal stigma but not stigmatisation, while 13.5% reported stigmatisation without personal stigma (66.1% report neither, 13.6% report both). This means that over one-third of respondents believe either that they should feel ashamed for claiming, or that other people think so.

[Table 3 about here]
There is similarly only a partial overlap between these measures and the take-up-related questions. People who agreed that a certain type of stigma existed were more likely to say that shame would affect their claim if they were needy and eligible (15.2, 3.6 and 8.5 percentage points more likely for each of personal stigma, stigmatisation and claims stigma respectively). Yet this still means that some people who disagreed that a certain type of stigma existed nevertheless said that shame would influence their claim (13.6% of those who disagreed there was personal stigma nevertheless said personal shame would influence their claim; this is 3.5% and 6.8% respectively for stigmatisation/social shaming and claims stigma/claims shaming). Stigma is therefore more widespread than any single question captures: 48.7% of respondents gave at least some sign of stigmatising benefits (personal stigma, stigmatisation, or a shame-related reason that would affect their decision to claim), and a further 30.2% disagreed that claimants are treated with respect.

**Stigma and ‘dependency culture’**

Finally, we test whether people in high-claim neighbourhoods hold lower benefits stigma than others; the results are shown in Table 4. (The table does not show results for claims stigma, which shows no significant patterns across any socioeconomic measure). In contrast to the ‘dependency culture’ hypothesis, the initial model shows that people in high-claim neighbourhoods report 5.4 percentage points more personal stigma than those in lower-claim neighbourhoods. When we simultaneously control for other socioeconomic factors then these patterns are maintained, and we also find a further surprising result: people in high-claim neighbourhoods are also 4.5 percentage points more likely to think that other people feel they should be ashamed to claim benefits (‘Stigmatisation’). Overall, those in high-claim areas were 4.6 percentage points more likely to give at least one stigma/shame-related response across the survey (‘All stigma/shame’).

Simultaneously, though, those in high-claim neighbourhoods were also less likely to say that personal shame would influence their decision to claim. This may reflect differences between morality and real-world preferences (as found for studies of the work ethic; Dunn, 2011:6), such that those in high-claim areas are more aware that need ultimately trumps stigma (Rogers-Dillon, 1995); or it may alternatively reflect differences in how stigma translates to shame, with those in high-claim neighbourhoods being more likely to contrast the deserving self with undeserving others (as discussed above). Table 4 similarly shows that other sociodemographic factors show reverse associations with stigma vs. shame as a reason for non-take-up, which may explain the inconsistent results found for the effect of education in the previous benefits stigma literature (Stuber and Schlesinger, 2006; Handler and Hollingsworth, 1969).
While not the focus of our study, it is worth noting that claimants themselves hold lower levels of personal stigma, but this is not the case for other forms of stigma/shame, and overall they are as likely to give a stigma-related response as non-claimants. There are also signs that out-of-work respondents and those in high-unemployment areas report some form of stigma more strongly (although these are not significant at conventional levels), and that those in social housing are less likely to give any stigma-related response. Many of the socioeconomic patterns are weak and/or non-significant, however, as has been found in smaller previous stigma surveys (Handler and Hollingsworth, 1969). Similarly, respondents aged 45+ were less likely to give a non-shame-related reason for non-claiming, but there were otherwise few significant patterns by age and gender.

[Table 4 about here]

Discussion

While stigma in the benefits system has long been debated, there have been few previous large-scale quantitative studies of benefits stigma, particularly in recent years. Using a new, nationally-representative survey with a boost sample of benefit claimants, we find that 10-12% of respondents agree themselves that people should feel ashamed to claim each of five individual benefits (‘personal stigma’) and 16-19% think that other people believe people should feel ashamed to claim (‘stigmatisation’). Looking at the proportions who report stigma for at least one benefit, 20% agree that personal stigma exists and 27% agree that stigmatisation exists – and as many as 34% say either one or the other. Moreover, an outright majority (58%) disagree that people are treated with respect when they claim (‘claims stigma’). It is therefore clear that benefits stigma is a multifaceted phenomenon, and that it is possible to capture this quantitatively.

We also asked respondents whether shame would influence their decision to claim benefits/tax credits, if they were needy and eligible. Over one-quarter (27%) say that a shame-related reason would make them less likely to claim; this is similar (if slightly higher) than given by eligible non-respondents as reasons for non-take-up in previous surveys. It should be noted, however, that we asked if shame would make people less likely to claim, and not if it would stop them claiming. Shame as an influence on take-up is also reported by some people who did not report personal stigma or stigmatisation, and in total, 49% of respondents give at least one stigma/shame-related response within the survey, with a further 30% disagreeing that claimants are treated with respect when applying. While the prevalence of some specific types of stigma for specific benefits can be relatively low, stigma in general is not a marginal phenomenon.
One of the strengths of our survey was that we could compare stigma between claimants and non-claimants, a comparison rarely available in the previous literature. Claimants of certain benefits (in-work tax credits, housing benefit, income support for single parents) are less likely to report personal stigma for that benefit. This may reflect that those who hold less stigmatising beliefs are more likely to claim benefits than others, or alternatively it may suggest a tendency for claimants to deflect stigma onto others (as shown in previous qualitative research). However, there are no differences for other types of benefits nor other measures of stigma, and overall, claimants are just as likely as non-claimants to provide at least one stigma-related response. While establishing a causal link between mental health and benefits stigma is problematic (the strongest study is Lee and Oguzoglu, 2007, who found weak, non-significant effects), the fact that one-third of claimants report personal stigma and/or stigmatisation, and 23% of claimants report a shame-related reason for delaying their claim in the past, is potentially concerning.

Finally, our study is the first to directly test whether benefits stigma is systematically lower in high-claim neighbourhoods, as the ‘dependency culture’ hypothesis predicts. Put simply, our results offer little support for this, fitting the results from qualitative research and survey studies of attitudes to work. Respondents in high-claim areas are less likely to say that personal shame would influence their decision to claim, but more likely to report personal stigma (and after adjustment, also stigmatisation). Overall, respondents in high-claim areas were more likely to report a stigma-related response, both before and after controlling for a variety of other socioeconomic factors. People’s decisions to claim benefits may be influenced by those around them, but this does not mean that claims are less stigmatised in high-claims area.

Recommendations for future research

While the Ipsos MORI benefit stigma survey has a number of strengths (e.g. operationalising sub-types of stigma, claimant boost samples) – and is available for other researchers to re-use – it also has several limitations. Some of these apply to all such surveys: for example, we are dependent upon self-reports of stigma, and these will be underestimates to the extent that it is stigmatising for someone to say they feel stigma. The quota sample may also have greater biases than a full random sample, although the careful design of the quota minimises this, and non-response biases will exist to varying degrees in all surveys.

Our experiences also suggest several improvements for future surveys. The prevalence of personal stigma and stigmatisation in this survey are lower than previous surveys, which may be because the present study asked about feeling ‘ashamed’, whereas previous studies asked about feelings that are both weaker and less stigmatising to admit (e.g. feeling ‘embarrassed’, ‘looked down on’, or a lack of ‘respect’). While this was a deliberate choice – shame has been the focus of recent scholarship and
is more policy-relevant – future surveys could capture both by allowing respondents to respond that (i) claimants should feel ashamed; (ii) claimants should feel embarrassed but not ashamed; or (iii) claimants should not feel embarrassed. It would also be helpful to make questions on claims stigma speak about ‘shame’ rather than ‘respect’, to make them comparable to questions on personal stigma/stigmatisation.

Our most surprising finding was the relatively small differences between the different benefits (much as these small differences were sometimes statistically significant), given that previous studies have fund considerable variation. One explanation is that the benefits involved – with the exception of tax credits (see below) – are similarly stigmatised because of their similar principles (predominantly needs-based benefits for the working-age population) and delivery (via DWP). Another explanation, however, is that many respondents interpreted the questions as about ‘deserving’ claims, and the stigma differences across these types of benefit are primarily about how far claimants are seen as deserving. One way of exploring this would be to ask explicitly about the stigma of deserving claims, and then to separately ask about the share of claimants that are seen to be deserving.

This raises a broader question: given the close link between stigma and deservingness, to what extent is it useful to study benefits stigma directly? Deservingness is a critical component of support for the benefits system (van Oorschot, 2000), and benefits policy therefore needs to carefully consider its impacts on deservingness perceptions (Baumberg, 2012). Nevertheless, our view is that there is some value in examining stigma and shame. Partly this is because shame is (by definition) a negative experience; even if many claimants deflect stigma onto others, shame is nonetheless a real aspect of a considerable minority of claimants’ experiences. Furthermore, stigma is part of the explanation – contentious in size, but not in existence – for the non-take-up of benefits by those who are in need and eligible, allowing hardships that the system was designed to avoid.

Policy implications
This paper has concentrated on describing benefits stigma, rather than in investigating how to reduce it. Nevertheless, other studies suggest three ways in which benefits stigma can be reduced. Firstly, political and media rhetoric could become less stigmatising. We have elsewhere examined the nature of media reporting of benefits in the UK, the role that politicians play within this, and how these might be changed (Baumberg et al., 2012). Secondly, claims stigma could be reduced by making the operation of the benefits system can be more respectful. We agree with e.g. the claimant-led Citizen's Commission on the Future of the Welfare State (1997:119) that benefits agency staff should be trained “to ensure that service users are treated positively, sympathetically and with respect, without stigma or hostility”, and that whenever claimants are asked to commit to
perform certain activities, personal advisors likewise sign a commitment to offer certain levels of support. Where claimants feel a greater entitlement to support, we would expect stigma to fall (Gubrium and Lødemel, 2014).

The third way of reducing stigma is perhaps the closest to the traditional heart of social policy: to change the structure of the benefits system. Britain has seen several deliberate attempts to create less-stigmatising benefits; 1960s research convinced Labour to replace the stigmatised ‘National Assistance’ with ‘Supplementary Benefits’ (according to Page, 1984:44), while tax credits in the 1990s were explicitly designed to – and appear to have succeeded in – lowering stigma (see also Breese, 2011). In general, we would expect less universal, less contributions-based, less generous, and more conditional benefits to be more stigmatised (Albrekt Larsen, 2006), with global studies of shame providing some support for this (Gubrium and Lødemel, 2014). As Richard Titmuss put it in Commitment to Welfare (1968:135), the ‘fundamental challenge’ of social policy is therefore to create a universal infrastructure around which selective provision can target services ‘with the minimum risk of stigma in favour of those who needs are greatest’.

However, this paper’s main contribution is descriptive rather than prescriptive: to show that sub-types of benefits stigma can be conceptualised and measured; that stigma is reported by a considerable minority of claimants and non-claimants alike; and that there is evidence against the idea that this overspills into a ‘dependency culture’. In this, it hopefully sheds new light on an old problem, and provides a further academic grounding for policy debates that seem likely to recur for as long as we have a benefits system in the present form.
Notes


3 Claimants will often be subject to other associated stigmas such as unemployment and poverty; but while these are a crucial part of lived experience, they are not part of our focus here.

4 Authors’ analysis of weighted data from British Social Attitudes survey 2000 (UK Data Archive SN4486).

5 For links to a number of headlines (and a critique of the statistics within them), see http://lartsocial.org/jameturnerstreet
References


Finn, D., Mason, D., Rahim, N. and Casebourne, J. (2008), Problems in the delivery of benefits, tax credits and employment services, York: Joseph Rowntree Foundation.


Stuber, J. and Schlesinger, M. (2006), 'Sources of stigma for means-tested government programs', Social Science & Medicine, 63, 933-945.


Taylor-Gooby, P. (2013), 'Why Do People Stigmatise the Poor at a Time of Rapidly Increasing Inequality, and What Can Be Done About It?', The Political Quarterly, 84: 1, 31-42.


### Table 1: Personal stigma and stigmatisation by benefit type (agree that should feel ashamed)

<table>
<thead>
<tr>
<th></th>
<th>All 1</th>
<th>Single parents</th>
<th>Unemployed</th>
<th>Incapacitated</th>
<th>Housing Benefit</th>
<th>Tax Credits</th>
</tr>
</thead>
<tbody>
<tr>
<td>Overall (self)</td>
<td>20.4%</td>
<td>12.1%</td>
<td>11.6%</td>
<td>10.3%</td>
<td>11.0%</td>
<td>10.1%</td>
</tr>
<tr>
<td>Claimants of that benefit</td>
<td>8.7%</td>
<td>5.8%</td>
<td>9.7%</td>
<td>10.4%</td>
<td>6.4%</td>
<td>6.8%</td>
</tr>
<tr>
<td>Overall (stigmatisation)</td>
<td>27.2%</td>
<td>19.1%</td>
<td>18.5%</td>
<td>17.0%</td>
<td>18.8%</td>
<td>16.0%</td>
</tr>
<tr>
<td>Claimants of that benefit</td>
<td>20.6%</td>
<td>18.3%</td>
<td>23.0%</td>
<td>22.3%</td>
<td>19.7%</td>
<td>18.6%</td>
</tr>
<tr>
<td>n (claimants)</td>
<td>936 1</td>
<td>132</td>
<td>233</td>
<td>273</td>
<td>454</td>
<td>290</td>
</tr>
</tbody>
</table>

Source: Ipsos MORI benefits stigma survey 2012; n(overall)=2478-2517. 1 For claimants, ‘all’=greatest stigma that a respondent reported about a benefit that they claimed.
Table 2: Stated reasons for not taking-up benefits/tax credits

<table>
<thead>
<tr>
<th>Type of benefit</th>
<th>Personal shaming</th>
<th>Social shaming</th>
<th>Claims shaming</th>
<th>Any shame</th>
<th>Non-shame</th>
<th>Would claim</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Benefits</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Overall</td>
<td>16.9%</td>
<td>4.3%</td>
<td>12.8%</td>
<td>26.6%</td>
<td>24.3%</td>
<td>56.6%</td>
</tr>
<tr>
<td>Claimants (past experiences)</td>
<td>14.2%</td>
<td>2.9%</td>
<td>11.3%</td>
<td>22.8%</td>
<td>25.2%</td>
<td>59.8%</td>
</tr>
<tr>
<td>Non-claimants (hypothetical)</td>
<td>17.6%</td>
<td>4.7%</td>
<td>13.2%</td>
<td>27.6%</td>
<td>24.0%</td>
<td>55.8%</td>
</tr>
<tr>
<td><strong>Tax credits</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Overall</td>
<td>13.0%</td>
<td>4.7%</td>
<td>12.9%</td>
<td>24.1%</td>
<td>25.2%</td>
<td>58.1%</td>
</tr>
<tr>
<td>Claimants (past experiences)</td>
<td>6.4%</td>
<td>2.1%</td>
<td>8.3%</td>
<td>14.1%</td>
<td>15.9%</td>
<td>74.9%</td>
</tr>
<tr>
<td>Non-claimants (hypothetical)</td>
<td>13.4%</td>
<td>4.9%</td>
<td>13.2%</td>
<td>24.7%</td>
<td>25.8%</td>
<td>57.0%</td>
</tr>
</tbody>
</table>

Benefits v. tax credits (sig.) 0.00* 0.52 0.82 0.02* 0.36 0.10

Source: Ipsos MORI benefits stigma survey 2012. *=p<0.05; n=2530.
### Table 3: Association between stigma per se and stigma as a reason for non-take-up

**Stated reasons for not taking-up benefits/tax credits**

<table>
<thead>
<tr>
<th>By levels of reported stigma</th>
<th>Personal shame</th>
<th>Social shaming</th>
<th>Claims shaming</th>
<th>Any shame-related</th>
<th>Non shame-related</th>
<th>Would claim</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>By felt stigma</strong></td>
<td><strong>ns</strong></td>
<td>ns</td>
<td>ns</td>
<td>ns</td>
<td>ns</td>
<td>ns</td>
</tr>
<tr>
<td>Among those disagreeing</td>
<td>13.6%</td>
<td>4.2%</td>
<td>13.2%</td>
<td>23.9%</td>
<td>23.8%</td>
<td>59.7%</td>
</tr>
<tr>
<td>Among neither disagree or agreeing</td>
<td>15.1%</td>
<td>2.8%</td>
<td>10.2%</td>
<td>23.9%</td>
<td>25.2%</td>
<td>56.3%</td>
</tr>
<tr>
<td>Among those agreeing</td>
<td>28.8%</td>
<td>5.6%</td>
<td>13.1%</td>
<td>36.8%</td>
<td>25.4%</td>
<td>47.1%</td>
</tr>
<tr>
<td><strong>By stigmatisation</strong></td>
<td><strong>ns</strong></td>
<td><strong>ns</strong></td>
<td><strong>ns</strong></td>
<td>ns</td>
<td>ns</td>
<td>ns</td>
</tr>
<tr>
<td>Among those disagreeing</td>
<td>14.4%</td>
<td>3.5%</td>
<td>11.6%</td>
<td>23.3%</td>
<td>24.7%</td>
<td>59.1%</td>
</tr>
<tr>
<td>Among neither disagree or agreeing</td>
<td>13.8%</td>
<td>2.8%</td>
<td>11.4%</td>
<td>24.4%</td>
<td>23.0%</td>
<td>58.4%</td>
</tr>
<tr>
<td>Among those agreeing</td>
<td>24.3%</td>
<td>7.1%</td>
<td>16.5%</td>
<td>34.8%</td>
<td>24.6%</td>
<td>50.4%</td>
</tr>
<tr>
<td><strong>By claims stigma</strong></td>
<td><strong>ns</strong></td>
<td>†</td>
<td><strong>ns</strong></td>
<td><strong>ns</strong></td>
<td><strong>ns</strong></td>
<td><strong>ns</strong></td>
</tr>
<tr>
<td>Among those disagreeing</td>
<td>16.0%</td>
<td>2.5%</td>
<td>6.8%</td>
<td>21.0%</td>
<td>19.8%</td>
<td>63.4%</td>
</tr>
<tr>
<td>Among neither disagree or agreeing</td>
<td>17.7%</td>
<td>5.1%</td>
<td>12.8%</td>
<td>28.6%</td>
<td>22.5%</td>
<td>56.3%</td>
</tr>
<tr>
<td>Among those agreeing</td>
<td>17.4%</td>
<td>4.6%</td>
<td>15.3%</td>
<td>28.5%</td>
<td>27.0%</td>
<td>53.5%</td>
</tr>
</tbody>
</table>

Source: Ipsos MORI benefits stigma survey 2012. Significance (for joint significance of differences between all three categories): † p<0.10, * p<0.05, ** p<0.01.
Table 4: Socio-demographic patterning of reported stigma

<table>
<thead>
<tr>
<th></th>
<th>All stigma/shame</th>
<th>Stigma per se</th>
<th>Reason for non-take-up</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Personal stigma</td>
<td>Stigmatisation</td>
</tr>
<tr>
<td>Neighbourhood claim rate</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Initial adjustment</td>
<td>3.7% +</td>
<td>5.4% **</td>
<td>3.3%</td>
</tr>
<tr>
<td>Adjustment for other factors</td>
<td>4.6% *</td>
<td>3.9% *</td>
<td>4.5% *</td>
</tr>
<tr>
<td>Other factors (mutually adjusted)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Personally claims out-of-work benefit</td>
<td>0.8%</td>
<td>-7.9% *</td>
<td>-3.5%</td>
</tr>
<tr>
<td>Local authority unemployment rate</td>
<td>5.0%</td>
<td>5.1% +</td>
<td>0.6%</td>
</tr>
<tr>
<td>Not working (vs. full-time worker)</td>
<td>4.1%</td>
<td>2.5%</td>
<td>5.4% +</td>
</tr>
<tr>
<td>Social housing (vs. owner-occupier)</td>
<td>-9.7% *</td>
<td>-4.3%</td>
<td>-6.3% +</td>
</tr>
<tr>
<td>No qualifications (vs. degree)</td>
<td>-5.1%</td>
<td>5.5%</td>
<td>-3.2%</td>
</tr>
</tbody>
</table>

Source: Ipsos MORI benefits stigma survey 2012. All estimates show average marginal percentage point effects; +=p<0.10, *=p<0.05, **=p<0.01. [1] 'Claim/unemployment rates' show the estimated difference between the middle of the top half vs. the middle of the bottom half of claims across areas (i.e. 75th vs. 25th percentiles). [2] 'Initial adjustment' is for age, gender, region, marital status, children, migration within the local authority, and personal benefit claims (out-of-work/tax credit/other). [3] 'Adjustment for other factors' additionally controls for local authority unemployment rate, work status, housing tenure, and qualifications.