The development and preliminary psychometric properties of two positive psychology outcome measures for people with dementia: the PPOM and the EID-Q

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Abstract

Background: Positive psychology research in dementia care has largely been confined to the qualitative literature because of the lack of robust outcome measures. The aim of this study was to develop positive psychology outcome measures for people with dementia.

Methods: Two measures were each developed in four stages. Firstly, literature reviews were conducted to identify and operationalise salient positive psychology themes in the qualitative literature and to examine existing measures of positive psychology. Secondly, themes were discussed within a qualitative study to add content validity for identified concepts \((n = 17)\). Thirdly, draft measures were submitted to a panel of experts for feedback \((n = 6)\). Finally, measures were used in a small-scale pilot study \((n = 33)\) to establish psychometric properties.

Results: Salient positive psychology themes were identified as hope, resilience, a sense of independence and social engagement. Existing measures of hope and resilience were adapted to form the Positive Psychology Outcome Measure (PPOM). Due to the inter-relatedness of independence and engagement for people with dementia, 28 items were developed for a new scale of Engagement and Independence in Dementia Questionnaire (EID-Q) following extensive qualitative work. Both measures demonstrated acceptable internal consistency \((\alpha = .849 \text{ and } \alpha = .907 \text{ respectively})\) and convergent validity.

Conclusions: Two new positive psychology outcome measures were developed using a robust four-stage procedure. Preliminary psychometric data was adequate and the measures were easy to use, and acceptable for people with dementia.

Keywords: Dementia, Positive psychology, Independence, Hope, Resilience, Outcome measure, Psychometric
and outcomes are under researched for people with dementia [6] and the reporting of positive experiences of people with dementia is limited to the qualitative literature.

To examine the positive psychology concepts and their relationship to wellbeing, valid and reliable outcome measures are needed. The aim of this study was to develop such measures to be used in further research and clinical practice. It is hoped this will lead to a more asset based approach to the study of dementia and enable people to further live well with their diagnosis.

Method

In order to develop these measures, we undertook a four stage process. Stage one consisted of a literature review in which evidence of positive psychology concepts from the qualitative dementia literature was sourced and examined. The most salient themes from these searches was developed into a topic guide for stage two. Stage two consisted of a qualitative study, in which positive psychology themes were explored with people with dementia and their carers to generate items for measures. Stage three consisted of expert feedback in which xdraft items within measures were examined for clarity, difficulty and content validity. Finally these measures were piloted with a small scale sample of people with dementia. These processes are discussed in more detail below.

Stage 1: literature review

Qualitative literature published between 1998 and 2015 that investigated positive psychology concepts for people with dementia was examined. Search terms were: goal, self-efficacy, hope, resilience, coping, wisdom, growth, sense of coherence, control, autonomy, pleasure, self-realisation, sense of agency, gratitude, happiness, optimism, transcendence, positive, dignity, social participation, social inclusion, self-concept, humour, creativity, flow, spirituality, love, compassion, benefit finding, community integration, opportunity, social adjustment, mindfulness, acceptance, successful aging AND dementia, lewy body, vascular, Alzheimer, cognitive impairment, old, elder. Truncations of search terms were used where appropriate. For the purpose of this section, the Seligman definition of positive psychology [7] was used to screen results. This search enabled the authors to identify salient concepts in order to produce a topic guide for the focus group stage of development. Concurrently, searches were conducted to identify existing, robust outcome measures that could be adapted for people with dementia. Search terms above were used again but search terms denoting dementia were changed for the search terms: measure, instrument, questionnaire, quiz, test.

Stage 2: focus groups

Design

A cross-sectional sectional qualitative design was used in which participants explored the meaning and implications of independence, social engagement, hope and resilience in dementia. Qualitative studies are recommended when designing outcome measures to ensure an adequate level of content validity [8]. A topic guide and semi-structured interview format were used to facilitate discussion and elicit in-depth views on subjective experience of this construct.

Participants

Ethical approval was obtained and participants were recruited through one private organisation (a care home in Leeds) and an existing Cognitive Stimulation Therapy (CST) group within a London NHS trust between September and October 2015. The CST group was identified as an appropriate source of recruitment for the focus group, as it was an established group consisting of people with mild dementia. People with dementia and carers were interviewed separately either individually or within a focus group, depending on preference and availability. 17 participants were recruited, all of whom met the inclusion criteria as detailed below:

Inclusion criteria

– People with a diagnosis of dementia according to the DSM-IV-TR criteria (American Psychiatric Association, 2000).
– Capacity to provide informed consent
– Able to communicate in English
  OR
– People who identified themselves as an informal carer to a person with dementia.
– Capacity to provide informed consent
– Able to communicate in English

Data collection

Interviews followed a semi-structured format that allowed interviewers to ask spontaneous questions that addressed individual circumstances. This ensured sensitivity to participants’ self-expression with regard to constructing his or her own accounts. Sessions were audio-recorded and discussion was facilitated around individual meanings and expectations of independence, social engagement, hope and resilience within dementia, the barriers and facilitators to these concepts and potentially related constructs including reciprocity. This enabled items to be developed that accounted for the individual differences of these concepts according to different participants.
People with dementia participated in two focus groups. However, due to the logistical difficulty in gathering members of carers in one space, individual interviews (some by telephone) were conducted with these participants. Examples of questions posed to both people with dementia and carers included: “Tell me about your (your relative/friend) experience of independence?” “How has independence changed for you (your relative/friend/person you care for) since you began having memory problems?” “How have social relationships changed for you (the person you care for)?” and “How does having people around you benefit or hinder you (the person you care for)?” Focus groups and interviews lasted between 16 min and 35 min and were largely dependent on the carer’s availability for individual interviews.

Data analysis
Thematic analyses are iterative in nature [9] and, to reflect this, transcripts were analysed by two researchers (CS and ML) independently. Initially, broad themes were generated to identify salient constructs. Following this, a consensus meeting was held in which researchers discussed their initial analyses. Any discrepancies between researchers were resolved through discussion. Once agreement was reached, both researchers’ broad themes were synthesised into a table and refined into codes. The primary author and an independent researcher (CS and AS) then reviewed codes and disagreements were discussed until a consensus was reached. To limit researcher bias, data was independently reviewed and analytical notes of researchers’ own inferences were recorded. In this way, potential biases are documented and can be explored if necessary [10].

Stage 3: expert feedback and revisions
Draft outcome measures were then submitted to a panel of experts, who were asked to provide feedback and pay particular attention to item clarity, difficulty and relevance. Experts were also asked to delete items they felt inappropriate and add any additional items to ensure the domain to be measured was captured in its entirety.

Stage 4: pilot study
Design
A cross-sectional design was employed in which participants completed two newly developed outcome measures, alongside three other outcome measures. Participants could complete the study in a number of ways depending on their preference. Firstly, in a self-report fashion, without supervision. Secondly, in a self-report fashion, supported by a member of the research team. Finally, in an interview led manner.

Participants
Participants were recruited primarily through existing CST groups within a London NHS Foundation Trust and via private and voluntary organisations (e.g. CogsClubs). Four CST groups were contacted and, of these four, two took part in the current study. Ethical approval was granted and the study was adopted for portfolio. The facilitators of these groups (e.g. facilitators of CST groups) identified participants who they deemed eligible to meet the inclusion criteria:

- Diagnosis of dementia according to DSM-IV criteria (American Psychiatric Association, 2000).
- Capacity to give informed consent.

Procedure
The researcher initially approached organisations detailing the study to determine whether there might be interest in participation. If group facilitators felt there might be interest, a date to attend the group was arranged. In cases where this was not possible, the study was introduced to the participants by facilitators after they had received all information regarding the study.

Participants were provided with an information sheet and a consent form prior to data collection and informal capacity assessments were conducted with all participants. Four people who were approached were deemed to lack capacity and therefore were not included within the current research. The majority of participants elected to have the researcher present (n = 26), hence, arrangements were made to visit participants at convenient dates and times. One CST group decided to complete the study during a break in a CST session (n = 4). For this group, the researcher was present to assist, if needed, as they completed the questionnaires. Three participants elected to take the questionnaire pack home with them and return it via post. These participants were provided with a pre-stamped envelope and the work mobile phone number of a member of the research team, should they encounter any difficulty in returning or completing the questionnaires.

Analysis
Initially, the homogeneity of domains was tested for both measures using an internal consistency analysis [11]. Following this, convergent validity was assessed using a Pearson’s correlation. Measures chosen to assess convergent validity were:

- Quality of Life in Alzheimer’s Disease Scale (QoL-AD) [12]
- Geriatric Depression Scale SF (GDS) [13].

The QoL-AD was selected as some people able to complete it satisfactorily with a MMSE score of three
decisions and advanced planning [23]. This relationship for tasks such as activities of daily living [22], financial 
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For example, people with dementia often require assist-
views on what it means to be truly independent. 
walk from one place to another. This may be a simplistic 
defined as a physical ability, for example, the time taken to 
mobility and decision making and this can be influenced by 
how others treat them. More recently, retention of inde-
tions of resilience of varying quality. Of these, the Connor-
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pendence has been investigated as a valuable outcome for 
the maintenance of wellbeing for people with dementia. It 
can decrease the potential stress felt by carers and delay 
nursing home entry [21]. However, independence is often 
defined as a physical ability, for example, the time taken to 
walk from one place to another. This may be a simplistic 
view, which does not take into account more subjective 
views on what it means to be truly independent. 

Independence may be closely related to social concepts 
including engaging with those around you and reciprocity. 
For example, people with dementia often require assist-
ance from their carer or immediate family to maintain a 
level of independence they are comfortable with. As de-
multiplicity and decision making and this can be observed 
would be between the PPOM, EID-Q and the QoL-AD. Positive psychology concepts may offer 
a protective effect against depression and, therefore, it was 
hypothesised that a negative correlation would be 
obscurred between the PPOM, EID-Q and the GDS.

Results
Stage 1: literature review
213 results were identified in which a number of positive concepts were explored. However, the majority of these papers were within the caregiving literature. Examples of qualitative studies reporting on people with dementia included a sense of coherence [17], spirituality [18] and hope [19]. A substantial review [20] of the qualitative literature indicated that resilience was particularly prominent within the qualitative literature, as did a qualitative study on the role of hope. Whilst independence and social engagement were discussed more sparingly and in the context of support or maintaining engagement with previously valued occupations, they may hold important implications for people with dementia. These concepts are discussed in more detail below.

Sense of independence and social engagement
The notion of a ‘sense of independence’ rarely appeared in the research literature. However, this may be particularly apt for people with dementia as they often face varying stages of independence within domains such as self-care, mobility and decision making and this can be influenced by how others treat them. More recently, retention of independence has been investigated as a valuable outcome for the maintenance of wellbeing for people with dementia. It can decrease the potential stress felt by carers and delay nursing home entry [21]. However, independence is often defined as a physical ability, for example, the time taken to walk from one place to another. This may be a simplistic view, which does not take into account more subjective views on what it means to be truly independent.

Independence may be closely related to social concepts including engaging with those around you and reciprocity. For example, people with dementia often require assistance from their carer or immediate family to maintain a level of independence they are comfortable with. As dementia progresses, a carer often assumes responsibility for tasks such as activities of daily living [22], financial decisions and advanced planning [23]. This relationship is of paramount importance with a supportive relationship being linked to slower cognitive decline for the person with dementia [24]. It is, therefore, reasonable to suggest that independence for people with dementia may be very closely related and dependent on this relationship. No suitable measures were identified that examined a sense of independence and social engagement, and this may be because previously independence was purely examined as a functional capability. Due to the relationship between these concepts, the decision was made to develop one outcome measure and items were based on discussions in the focus group study.

Resilience
Resilience is often used to describe those who display emotional stamina in adverse situations. Its definition is sometimes ambiguous and therefore, the definition used in the current study will be the flexibility in response to changing situational demands [25]. It is noted that whilst some definitions refer explicitly to an adverse event preceding the development of resilience, the definition presented refers to changing situational demands as the precedent for resilience rather than an overtly negative event. Whilst it can be suggested that resilience is not an entirely positive construct, it is included in the current research due to its presence in a recent systematic review of living positively with dementia [26]. More specifically, this review of qualitative literature identified themes of engaging with dementia and ‘facing it and fighting it’ as a form of active perseverance through resilience. An earlier systematic review [26] identified four measures of resilience of varying quality. Of these, the Connor-Davidson Resilience Scale (CD-RISC) [27] was selected for adaption due to its thorough assessment of sensitivity to change and interpretability. However, the CD-RISC is a 25-item measure and, it was decided that this could be too time intensive. As such, the short form version (CD-RISD-SF) was examined and found to have adequate psychometric properties [28].

Hope
A substantial qualitative study examined the presence and conceptualisation of hope for people with dementia, for which their findings provide a strong rationale for the applicability of multidimensional hope, rather than goal directed hope [19]. For most, hope was described as an active process whereby a developmental history of ‘learned hope’ remains well preserved and this facilitated a process of ‘keeping going,’ viewing difficulties encountered as challenges to overcome. Furthermore, there appeared to be an active re-appraisal of balancing hope and realism. This seemed to indicate that for people with dementia, outcomes hoped for are grounded in realism and generalised in nature, as illustrated by themes of
themes are discussed briefly below with selected quotes and its relatedness to sense of independence. These actions with those around them and illustrates the desire of people with dementia to engage in social interaction also displayed a retained desire and satisfaction in independence and this was often discussed by carers in the context of isolation from others: ‘I think it’s being able to do things by yourself maybe without the assistance of others or just being free to choose’ (C1). However, when asked to explore what constituted independence for people with dementia, definitions became more varied. Most prominently, participants felt that independence changed as ability to complete activities of daily living declined. This led to interdependence between a person with dementia and a carer. Examples of what this constituted appeared to be the giving of slight assistance to people with dementia so that they could maintain a level of independence that suited their individual ability: ‘Dressing and an occasional prompt... when the automatic is no longer automatic’ (C4).

**Stage 2: focus group**
As no measures of sense of independence and related social concepts were identified within stage 1, specific attention was paid to these concepts during focus groups and interviews. The total sample was 17, in which two focus groups for people with dementia were used (Group 1 n = 3; Group 2 n = 6) and individual interviews were employed for carers (Table 1).

Four overarching themes emerged as central to independence in dementia: 1) independence and interdependence, 2) functional independence, 3) remaining active and 4) social engagement. The first higher order theme illustrates ambiguity in definitions of independence in dementia and indicates that a period of interdependence between a carer and people with dementia can be beneficial for both. The second and third higher order themes reflect the differing domains within the construct of ‘independence’ and suggest that physical and mental ability may have important implications on the retention of independence. This was often compensated for with highly individualistic support provided by carers and a constant reappraisal of abilities and tasks as people with dementia’s ability to engage in such activities declined. The final higher order theme describes the retained desire of people with dementia to engage in social interaction with those around them and illustrates the barriers and facilitators to maintaining this engagement and its relatedness to sense of independence. These themes are discussed briefly below with selected quotes to support their applicability.

### Table 1 Participant demographics

|                        | Age M (SD) | Female n (%) | Ethnicity: Caring for: | Caring for: |
|------------------------|------------|--------------|------------------------|-------------|
| PwDa n = 9             | 80.56 (±7.13) | 7 (77.8%) | Asian: 1 | n/a |
| White: 8               |            |              |                        |             |
| Carer n = 8            | n/a        | 0.8          | Asian: 1 | Parent: 3 |
| White: 7 | Spouse: 4 | Grandparent: 1 |                      |             |
| TOTAL n = 17           |            |              |                        |             |

*Person with Dementia (PwD)*

**Independence and interdependence: definitions outside of and within dementia**
Participants were initially asked to explore definitions of independence and this was often discussed by carers in the context of isolation from others: ‘I think it’s being able to do things by yourself maybe without the assistance of others or just being free to choose’ (C1). However, when asked to explore what constituted independence for people with dementia, definitions became more varied. Most prominently, participants felt that independence changed as ability to complete activities of daily living declined. This led to interdependence between a person with dementia and a carer. Examples of what this constituted appeared to be the giving of slight assistance to people with dementia so that they could maintain a level of independence that suited their individual ability: ‘Dressing and an occasional prompt... when the automatic is no longer automatic’ (C4).

**Functional independence: activities of daily living, self-care and decision-making**
Independence was discussed within the context of giving the ‘right’ level of support for a person with dementia. Too much could be frustrating for the person with dementia and too little could lead to the person with dementia feeling neglected. Carers often felt that, by adjusting the level of support given to a person with dementia based on their ability, it allowed people with dementia to focus on other decisions: ‘yeah just having sort of basic needs assisted with and then you’ve got extra energy to focus on some other things that might be a bit more challenging’ (C8). The potential for conflict occurred when the person with dementia felt that aspects of self-care were being taken out of their control but still appeared desirous to make a decision even if this was, as seen by the carer, the wrong decision: ‘If I say I don’t think you should... wear that... It’s not suitable for the weather. No he wants to wear it... if I don’t let go he’s very grumpy, upset’ (C2). People with dementia also displayed a retained desire and satisfaction in decision-making: ‘because I like to make decisions for myself...’ (P1)

**Remaining active**
Physical ability was implicitly related to maintaining a level of independence: ‘because I don’t go out... my legs. I like to go out and get on the bus to Romford... I couldn’t walk very far that’s the trouble with my legs’ (P5). As physical ability decreased, complex hobbies were often abandoned ‘I got plenty (of hobbies) but I can’t do them now’ (P3) and this could impact upon their sense of independence. An example of this came from a participant who discussed being no longer able to fish: ‘I’ve got to roll myself over to a spot so that I can
just get up and not join the fish... it's impossible' (P8), suggesting a close relationship between physical abilities and remaining active, and its impact on ability to remain independent. For carers, the need for people with dementia to engage in activities was seen as a form of mental stimulation, to delay dependence for as long as possible: 'It must keep the bits of her brain that are still functioning more active than they are currently and that just cannot be a bad thing' (C3).

**Social engagement**
Social concepts were often discussed in close relation with independence. For example: 'Being able to make decisions yourself without bothering anyone else but at the same time taking their problems into concern' (P1) and independent social engagement allowed people to feel they had accomplished much: 'we often say oh I'll pop out for an hour or so tonight...you feel fulfilled then at the end of the day and you don't need much if you're not talking about anything much' (P2). It was noted that people with dementia wanted varying levels of engagement and independently choosing their own level of engagement appeared to be the most beneficial: 'He was quite happy in a large pub where he could wander around and you know just chose who he would talk' (C1).

Reciprocity was seen by both people with dementia and carers as a means of being useful to a carer in ways that were manageable and was always discussed as an exchange of either activities or sentiment: 'Being kind to one another' (P1). A desire to give back was also explored by carers who were aware of the person with dementia's desire to be reciprocal with them: 'He knows that it would please me and it would make me happier if he helped. This is why he does it. I wouldn't say it's his hobby' (C2). However, when a person with dementia felt they could not be reciprocal, it could result in feelings of guilt and frustration: 'I mean bless his heart he says to me now I've ruined your life... he gets cheesed off at the fact his daughter is doing what I'm doing' (C5). This latter example was discussed within the context of a carer doing home improvement tasks around the house; something the person with dementia had been responsible for prior to the onset of dementia. This loss in autonomy resulted in friction within the relationship between the carer and person with dementia.

**Resilience and hope**
Saturation for resilience and hope data was reached very quickly. Resilience appeared to be present for people with dementia and was defined as: ‘Take it as it comes unless you really clever and want to do something but I'm not’ (P2). This attitude of taking life as it comes was discussed with regard to various hospital appointments and health problems. As such, health related resilience might hold more importance than other aspects of resilience for this population and people with dementia noted the additional implications of ageing on health: ‘The older you get the more difficult it becomes’ (P1). In line with previous work, people with dementia viewed hope generally rather than situation specific and as being present on a day-to-day basis: 'Well I say I hope I get this and I hope I get that' (P1). The applicability of future oriented hope was also examined by one participant: 'I feel scared about my future but maybe I don't know if it's further down but just on a day-to-day you are... just worried about the day' (C2). This may be particularly relevant for people with dementia, as participants with dementia did not discuss future oriented hope.

**The EID-Q and PPOM**
Using the results of stage 1 and 2, 28-items were generated for the Engagement and Independence in Dementia Questionnaire (EID-Q). Due to the inter-relatedness of independence and social engagement, the decision was made to combine these concepts within one scale. As hope and resilience are more representative of positive psychology traditionally, The Herth Hope Index and CD-RISC-SF were combined and item wording was adapted to create the Positive Psychology Outcome Measure (PPOM).

**Stage 3: expert feedback and revisions**
Five experts from the Promoting Independence in Dementia (PRIDE) research programme responded to a request for feedback and revisions. Professions consisted of a professor of old age psychiatry, a reader in clinical psychology, a professor of community care research, a psychologist and a researcher. Results from all responders were pooled into one document using track changes and discussed within the supervisory team, until a consensus as to which suggestions to include as amendments was met. Most suggestions were acted upon to ensure an adequate level of content validity.

**EID-Q**
Responders noted three items that should be removed, as they didn't appear closely related to the concept in question. Examples of items removed included 'I feel connected to society'. It was suggested that other items were in need of rewording as they may be too difficult to understand. Six items were reworded to improve clarity. Examples included 'I can adapt my wishes to be in line with what I can do,' which was replaced by 'I can make changes to my life to match my ability' and 'I can participate in a meaningful conversation,' which was amended to 'I can participate in discussions in ways that I enjoy'. No additional items were suggested but one responder did comment that the overall length might be excessive.
Responders did not identify any redundant items within the PPOM but suggested that a number of items needed rewording to make them more appropriate for people with dementia. 9 items were reworded and examples included rewording ‘I can see possibilities in the midst of difficulties’ to ‘I can see positive things in difficult situations’ and ‘I am able to adapt’ to ‘I am able to adapt to things’. Other items were suggested as in need of clarifying. For example, ‘I am a strong person’ was amended to ‘I am an emotionally strong person’.

Stage 4 pilot testing

Thirty-eight people with dementia were approached to complete the novel measures and convergent validity measures. Of these, four lacked the capacity to consent and one declined to take part. This left a sample size of 33 (Table 2). All participants had been diagnosed with dementia and were deemed capable of giving informed consent. Eight participants elected to complete the measures in an interview led manner, whilst 25 elected to complete it in a self-report manner. All data was inputted into an SPSS file on a password-protected computer. Reverse items were recoded on the PPOM and EID-Q so that higher scores indicated a higher level of the domains assessed.

Internal consistency

PPOM The first internal consistency analysis revealed that the PPOM had an overall Cronbach alpha level of $\alpha = .793$, and the subscales were $\alpha = .859$ for resilience and $\alpha = .557$ for hope. This highlighted that one or more of the items on the hope subscale may not be pertinent for people with dementia. The resilience subscales’ internal consistency was almost identical to the original, reported as $\alpha = 0.85$ (Campbell-Sills & Stein, 2007) [24]. Following further analysis, it emerged that the removal of three items on the hope (‘I feel all alone’; ‘I have faith in the future’; ‘I feel scared about the future’;) subscale and two items on the resilience subscale (‘I can achieve my goals’; ‘I am not easily discouraged’) would improve the internal consistency of the PPOM. After removing these items, the overall internal consistency improved to $\alpha = .849$, with the subscales improving to $\alpha = .755$ for hope and $\alpha = .871$ for resilience. This therefore resulted in a 16 item scale, consisting of a hope subscale (8 item) and resilience subscale (8 item).

EID-Q Initially the EID-Q had an adequate level of internal consistency ($\alpha = .896$) and the subscales were also of sufficient value ($\alpha = .849$ for independence and $\alpha = .771$ for engagement). However, by removing one item from each subscale (‘there are things I would like to do but can’t; I have good relationships/friendships with others’) internal consistency was raised to $\alpha = .868$ for independence, $\alpha = .775$ for social engagement and $\alpha = .907$ for the scale overall. This resulted in a 26 item scale, consisting of an independence subscale (13 item) and engagement subscale (13 item).
Convergent validity

PPOM Preliminary indications of convergent validity were found between the PPOM and the GDS. A two-tailed Pearson’s R correlation found a significant, negative relationship between the PPOM and the GDS (r = -0.562, p = .002) and the hope subscale and the GDS (r = -0.557, p = .001). However, the resilience subscale did not significantly correlate with GDS (r = -0.312, p = 0.088). An additional Pearson’s R correlation was performed to assess the relationship between scores on the QoL-AD and the PPOM but no significant correlations were found.

EID-Q As with the PPOM, a two-tailed Pearson’s R correlation was performed for both subscales and the EID-Q scale and the GDS. Significant correlations were found between the independence subscale (r = -0.447, p = .012) and GDS, the engagement subscale (r = -0.430, p = .016) and the GDS and the EID-Q and the GDS (r = -0.461, p = .009). This again indicates a negative correlation between the sense of independence, social engagement and depression. Pearson’s R correlations between the EID-Q and the QoL-AD indicated an emerging positive relationship between independence, engagement and quality of life. Firstly, the total QoL-AD score was found to be positively correlated with the independence subscale (r = 0.497, p = .005), engagement was correlated with the QoL-AD (r = 0.586, p = .001) and the EID-Q was found to be positively correlated with the total QoL-AD score (r = 0.557, p = .001) (Table 3).

Discussion
Summary of results
This is, to our knowledge, the first study to report on the development of positive psychology outcome measures for people with dementia. The Positive Psychology Outcome Measure (PPOM) and the Engagement and Independence in Dementia Questionnaire (EID-Q) were developed using a robust four-stage procedure resulting in robust positive psychology outcome measures (Additional file 1). Potentially relevant concepts and related outcome measures were sourced and examined in the current literature. Content validity was provided during focus group studies and expert revision. Finally, a small-scale pilot established adequate psychometric data for the new measures. The two new measures were easy to use and acceptable to participants, with most being able to give an informed opinion as to the use of these concepts in everyday life.

Findings in the context of the literature
Increasingly, the aim of psychosocial research for people with dementia is to maintain independence through interventions including exercise [33] and occupational therapy [34]. However, within these studies independence is defined in varying forms and usually as a functional ability, for example, the time taken to walk from one place to another or the ability to dress, both without the assistance from others. Whilst there appears to be an implicit expectation that improving independence will increase wellbeing, the observed relationship can be ambiguous [35]. The current study used definitions of independence as discussed by people with dementia in a qualitative setting and as such, the EID-Q is a more holistic measure of a sense of independence, rather than a measure of an operant capability.

The importance of the relationship between a person with dementia and their carer has previously been explored with a reciprocal relationship being highlighted as an important feature [36] and it has been proposed that reciprocity is a potential means of mitigating a loss of autonomy [37]. This reciprocal relationship refers to the ability of the person with dementia to provide assistance to their carer in ways that they feel are beneficial to them, in exchange for the carer providing means of assistance to them. Reciprocal relationships have been found to have numerous health benefits including reducing risk of stroke [38] and increasing psychological wellbeing for older adults [39].

Accounts of hope within the current study were consistent with previous findings in which future oriented hope appeared to be less appropriate for people with dementia [40] and accounts of resilience appeared to support the definition of adapting to changing situational and personal demands. Evidence for the relationship between hope and depression was also documented here and is consistent with the theory that positive psychology concepts may protect against depression [41, 42]. However no significant correlation was found between the PPOM and QoL-AD, possibly indicating hope and resilience may be distinct from quality of life.

Resilience as a positive psychology trait is not without issue. It is generally accepted that resilience is not a construct entirely characteristic of positive psychology but it was included within the current research due to its presence in the largest qualitative study of positive psychology and people with dementia to date [20]. Furthermore, positive psychology seeks to provide a balance and to exclude such a salient

| Table 3 Pearson’s R correlations between measures | GDS | QoL-AD |
|-----------------------------------------------|-----|--------|
| PPOM Total: | r = -0.562, p = .02 | - |
| PPOM: Hope | r = -0.557, p = .001 | - |
| PPOM: Resilience | r = -0.312, p = .088 | - |
| EID-Q Total | r = -0.461, p = .009 | r = 0.557, p = .001 |
| EID-Q: Independence | r = -0.447, p = .012 | r = 0.497, p = .005 |
| EID-Q: Engagement | r = -0.430, p = .016 | r = 0.586, p = .001 |
characteristic could leave to a tyranny of positives in which potentially beneficial concepts for wellbeing are not examined, because it is not seen as a completely positive trait.

Currently, there is a wealth of positive psychology outcome measures in use for other populations [43] but this approach is only just beginning to be applied to those with dementia research. There is an increasing recognition that positive factors can act as a protective agent in the development of negative emotional states including depression, and this has been exemplified in the dementia caregiving literature [44], where positive aspects of caregiving has been proposed as a buffer the negative consequences of caregiving such as depression, burnout and medical risk. The positive psychology approach to dementia reflects an asset-based standpoint, in which the retained capabilities and strengths that people with dementia exhibit are examined. These retained capabilities or strengths may be an important contributing factor to a person’s wellbeing and interventions could be designed to bolster or maintain these strengths over time.

Methodological problems and limitations
Preliminary psychometric properties were established with a relatively small sample size, increasing the risk of a type II error. Also, the format in which measures were delivered may have impacted upon the instances of missing data. 25 participants elected to complete the measures in a self-report manner and these participants were more likely to miss items than those who completed the study in an interview led manner. This may have been a formatting issue or that participants felt unable to make judgments on particular items without guidance. Certainly within the interview led sample, participants struggled to answer future oriented questions within the hope subscale for the PPOM. This is supported by previous research in which goal oriented hope was found to be lacking for people with dementia [19]. Future oriented questions regarding hope were subsequently removed from the measure. Furthermore, it was noted that carers often had differing perspectives to people with dementia. This was particularly evident for the item ‘I can look after myself as much as I need to’ and it is unclear how much this affected the self-report sample.

We did not collect information with regard to cognition as the qualitative literature suggests that people with dementia experience these positive emotions throughout the course of dementia [20] and as such scores on a cognition test would not be relevant to these measures. However, all participants here had capacity to provide informed consent and so it can be assumed they were in the more mild stages of dementia. We therefore cannot make assumptions about the experience of positive emotions in later stages of dementia. This is something future researchers may wish to examine further.

Finally, it must be noted that a proportion of our participants were also receiving interventions including Cognitive Stimulation Therapy. These participants may differ slightly to those not engaging in activities but this is mitigated by the pilot study sample, for which participants who were not receiving an intervention were also involved. This limitation will be addressed in a further larger scale study.

Future research
As this study only provided tentative psychometric properties for the PPOM and EID-Q, the next stage is a larger study and further psychometric analysis, which is currently ongoing, and will allow the testing of the measures test-retest reliability, discriminant validity, factor structure and responsiveness. This will ensure that the developed measures can be applied within dementia services and research across the country.

Positive psychology is a broad discipline and there appears to be a need to formulate a theoretical model of this approach tailored specifically for dementia, to recognise the unique difficulties and capabilities this population has. This has, to some extent, been established within the caregiving literature with a focus on positive experiences of caregiving [45, 46] but is currently lacking within the dementia literature.

The evaluation of positive psychology concepts share difficulties with the appraisal of quality of life as it is argued that, below a certain level of function, people with dementia struggle to accurately appraise this [47]. Nevertheless, this study illustrates that, whilst positive psychology concepts require complicated appraisals, people with dementia appear able to give a view to this. As such, future quantitative research is needed to examine other concepts that may hold significance for people with dementia including love and humour.

Conclusion
Two positive psychology measures were developed using a robust, iterative process. The EID-Q and PPOM demonstrated acceptable internal consistency and convergent validity. However these psychometric properties were established using a small sample. Further psychometric analysis is needed before such measures can be used in psychosocial research to improve our knowledge and understanding of wellbeing for people with dementia.

Additional file

Additional file 1: The Engagement and Independence in Dementia Questionnaire (EIDQ) and Positive Psychology Outcome Measure (PPOM). Description of Data: The EID-Q (26-item) and the PPOM (16-item). Both are answered on a 5-point Likert scale, with a timescale of the previous month. (DOCX 107 kb)
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Availability of data and materials
The datasets used and/or analysed during the current study available from the corresponding author on reasonable request.

Authors contributions
CS undertook data collection, data analysis and writing of the manuscript. MD commented on drafts of the paper, provided methodological support and supervised the study. ML undertook a section of data analysis and commented on drafts of the manuscript. EC commented on drafts of the paper and provided supervisory support. AS designed the study in collaboration with CS. AS also commented on drafts of the manuscript and provided supervisory support. All authors read and approved the final manuscript.

Competing interests
The authors declare that they have no competing interests.

Consent for publication
Not applicable.

Ethics approval and consent to participate
Ethical approval for the current study was granted in two parts. Firstly, as part of the Promoting Independence in Dementia research programme (15/EE/0134) at the East of England Research Ethics Committee (REC) and secondly as part of an individual PhD study (15/EE/0443), again at the East of England REC. All participants who participated were deemed capable of providing informed consent and were required to sign a consent form.

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