**Maximising engagement and participation of intellectual disability (ID) staff in research: insights from conducting a UK-wide survey**

**Authors**

**Claire Kar Kei Lam** *Kingston University & St George’s, University of London*

**Jane Bernal** *University of South Wales*

**Janet Finlayson** *Glasgow Caledonian University*

**Stuart Todd** *University of South Wales*

**Laurence Taggart** *University of Ulster*

**Annette Boaz** *Kingston University & St George’s, University of London*

**Irene Tuffrey-Wijne** *Kingston University & St George’s, University of London*

**Abstract**

**Aim:** This paper explores ways of maximising research engagement of ID staff as research participants, research advisors and research implementors.

**Method:** The authors describe and reflect on a three-phased strategy in recruiting frontline staff (n=690) working for ID service providers (n=25) to participate in a UK-wide anonymous online survey about death, dying and bereavement.

**Results:** Important elements in engaging participants were: involving stakeholders at all stages of the research process, which includes building relationships with participating organisations; enlisting organisational management support at all levels; an attractive and well laid-out collection tool; a well-structured recruitment strategy; time and flexibility; and a varied and targeted dissemination strategy. However, the recruitment method had limitations, in particular around representativeness, bias and generalisability.

**Conclusions:** Staff in ID services can be enthusiastic and invaluable research participants. Active engagement between researchers, participating organisations and stakeholder groups is key to ensuring involvement of ID staff with research.

**Key words**

Difficult topic, Intellectual disability staff, Research methods, Response rates, Surveys

 **Background**

This paper focuses on ways of maximising research engagement of staff working in intellectual disabilities (ID) services, mostly in terms of recruiting ID staff as research participants, but also in terms of their contributions as research advisors and research implementers. Active engagement of staff with research is one way of bridging the gap between research and practice. This, in turn, can facilitate changes in staff attitudes and in the way things are done; promote improved health and well-being of staff and clients; and influence policy changes (Read, 2018). Achieving this can be especially challenging when the research is concerned with a difficult topic, or even a taboo topic; we draw on our experience of doing research around dying, death and ID.

In particular, this paper reflects on the methods, processes and outcomes of a UK-wide survey of 690 staff working with people with ID in residential and supported living services, and how these research processes affected stakeholder engagement with the study. The *Talking About Dying Survey* investigated how the staff communicate about death and dying with people with ID who are dying or bereaved; how they confront issues of death, dying, and bereavement at work; and how often they were confronted with such issues. The survey was developed and conducted in 2017-2018 by the authors, a team of collaborating researchers based at four universities across all four UK countries.

There have been many studies where ID staff took part as research participants and informants. For example, with regards to our own research topic, a systematic literature review on the experiences of staff who support people with intellectual disabilities around death, dying and bereavement found 13 papers reporting studies where frontline care staff and/or managers were the participants (Lord *et al.*, 2017). Eleven of these were qualitative studies involving focus groups or interviews, one was a qualitative questionnaire (n=38) (Forrester-Jones, 2013), and one was a survey with qualitative responses (n=57) (Hoover *et al.*, 2005). These staff-focused studies highlight the importance of investigating staff perceptions, knowledge and attitudes, as appropriate support for people with intellectual disabilities around death and dying depends on staff who feel confident and are well supported themselves.

A Dutch survey of staff working directly with people with ID who have end of life care needs (Bekkema *et al.*, 2015) included 294 questionnaires sent to care staff employed by care services for people with ID; of these, 196 (67%) were returned. These staff were more highly educated than most frontline ID support staff in the UK: 85 were registered nurses, 8 were certified nursing assistants and 103 were social workers; all were members of a national research panel, which may explain the good response rate.

Hunt et al. (2019) report on a survey investigating end of life care outcomes for people with ID living in residential care in the UK. They sent a detailed post-bereavement questionnaire to 188 ID staff involved in the support of a person with ID who was known to have died, of which 158 were returned. Core details about 222 deaths, and the staff contact details for 188 of these, had been obtained from the ID service providers.

There is a dearth of literature around the methodological issues of involving ID staff as research participants and informants. The published papers indicate that at least some ID staff are willing to participate in research on difficult topics, but they do not provide much detail on the design and conduct of death-related studies within ID services; nor do they shine a light on how to engage staff with the topic. There is little evidence detailing the process of recruiting care staff, outlining any pitfalls when involving support staff in ID research, or outlining recommendations for successful engagement and satisfactory levels of participant recruitment.

Hall et al. (2017) conducted a small focus group study of ID staff, aimed at identifying ways to maximise recruitment of study participants with ID, facilitated by ID staff. They made three suggestions for researchers, which may also be of relevance for studies where the staff themselves are the main participants: (1) flexible contact methods, aided by making use of digital avenues; (2) forming stronger relationships between universities and care companies, which can help disseminate research findings, and develop continued collaborations in future studies; and (3) approaching managers as the main contact, as supportive managers are the key factor to staff recruitment. This confirms a suggestion by Tuffrey-Wijne et al. (2008) that ID staff are more likely to participate (or facilitate participation of people with ID) if they can see personal benefit, such as useful study outcomes or the cathartic effect for the study participant of engaging with researchers on a sensitive topic.

Our objectives for this paper are to describe our strategy in recruiting ID support staff to participate in our survey, and to explore and evaluate the factors that affected the engagement between researchers and practitioners, including the response rate and the engagement of ID managers and frontline staff with the disseminated findings. We will also explore the limitations to our methods, some of which were significant. We hope this will be of use to anyone planning to conduct research involving ID staff and services, not just in the UK but internationally.

**Methods**

***The Talking About Dying Survey (TADS)***

The TADS questions were based on the research evidence of current issues and best practice with regards to death-related communication (Ryan *et al.*, 2011; Todd, 2013; Tuffrey-Wijne, 2013; Wiese *et al.*, 2013; Stancliffe *et al.*, 2016; Tuffey-Wijne and Rose, 2017; Northway *et al.*, 2018). The study objectives included gauging how current practice relates to this evidence base, as well as adding to the evidence by gathering new data on staff issues and concerns. This, in turn, would inform the development of guidance and training. We received 690 completed online questionnaires from staff working for ID providers of residential and supported living services across the UK (61% response rate). Of these respondents, 68% were direct support staff, 29% were frontline managers, and 3% were “other”. The study findings are reported elsewhere (Tuffrey-Wijne *et al*., 2020).

***Research Advisory Group (RAG)***

The RAG comprised of 16 members lending different expertise and perspectives to the TADS research team, including three people with ID, two family carers of people with ID who had been bereaved or had died, two senior managers from ID service providers, an ID nurse, two ID support staff, and four academic researchers. Several of these stakeholders were involved from the time of developing the study protocol. The RAG met four times throughout the project, from before the design of the questionnaire through to the dissemination of findings towards the end of the project. This stakeholder involvement ensured that the research questions and study design were relevant to its target audience. For example, a family carer pointed out the importance of asking questions about people’s understanding of death. Her adult son with Down Syndrome knew that his father had died, but had difficulty understanding that he would never come back, making it more difficult for the mother to talk about the death with her son. As a result, we included two questions related to the person’s understanding of the universality and permanence of death (“everybody dies” and “people who die can never come back”). Senior managers from ID service providers held additional conference calls with the research team to discuss participant recruitment strategies, including the feasibility for their staff to complete a 20-40 minute survey online.

***Survey design***

This was an anonymous questionnaire, using an online survey tool (SurveyMonkey©). Staff who had experience of supporting a person with ID in their care who had died, was terminally ill, or had been bereaved in the past year, were asked detailed questions about this person and about their own experience of supporting them. On a topic as difficult as death and dying, great sensitivity was needed. It was essential that the survey was well presented and engaging, and that respondents felt it was relevant to them. All RAG members helped trial the survey by completing it themselves and sharing it with peers; they then met together to share feedback. This enabled us to assess how long it took to complete the survey and where the risk of respondents abandoning the survey was the greatest. At these risk points, we inserted photographs and graphics, for example, a photo of two research advisors with ID holding up encouraging signs saying *Please keep going! It will really help to know more about this.* Edits were made to clarify, rephrase, and replace unnecessary or unclear questions. This was helped by using the software’s internal logic tool, which ensured that questions that were not relevant to a respondent were skipped automatically. We ensured that respondents’ answers would be saved automatically as long as they returned to the same device and browser; this allowed respondents to quit and return to the survey at any time, which was useful as staff often had several other demands on their time. The survey was trialled in all available formats (tablets, mobile phones, and desktop computers) with 39 stakeholders and adjusted in light of their feedback, before inviting participants.

***Three-phase participant recruitment***

The targeted survey respondents were support staff working directly with people with ID, either with or without recent experience of death or bereavement among the people they supported. We utilised a top-down recruitment approach in three separate phases: (I) recruiting organisations; (II) recruiting middle managers; (III) recruiting survey respondents (direct care staff) [see Figure 1].

We decided that only one staff member per team or residence should be invited to complete the survey. This was not only to ensure that the burden on staff teams was not too great, but also to ensure that there would not be more than one survey response relating to a particular deceased or bereaved person with ID, as this would skew the findings in relation to the profiles of people with ID who have died or are bereaved. The survey was anonymous, partly to encourage honest responses, and partly to comply with the demands of theKingstonUniversity Faculty Research Ethics Committee (REC) which approved the study. The REC prevented us from contacting any respondents directly, in contrast to the survey by Hunt et al (2019), where ID staff who had supported a deceased person with ID were identified by the research team and non-responders could be followed up. Our gatekeepers were ‘middle managers’, i.e. those with direct managerial responsibility for a range of staff teams, whom we relied on to select potential respondents, help us relay information to their staff such as the survey link, reminders, and information sheets to the intended participants.

The anonymous nature of the survey probably helped respondents to give honest responses, but it also had drawbacks. We had no way of identifying (for example by initials, date of birth, or location) the people with ID whose death or bereavements staff were describing. The approving REC ruled that ethical considerations also prevented us from allocating a unique identifying number to each care setting, because it might have enabled either the staff member or the person with ID to be identified. There was therefore no way to detect when more than one staff member from a single setting had sent in a response, other than the original direction to middle managers to send the survey to only a single member of support staff in each setting. We have some evidence of one likely instance of double counting, where several respondents described a person with ID who had died of an extremely rare condition, with similar other characteristics, in the same county. The inability to detect double counting was a potential flaw in this methodology. It negatively affecting our confidence in some of the findings, in particular those findings relating to the profiles of deceased or bereaved people with ID.

*Phase I: Recruiting organisations*

A total of 59 ID service providers within the UK were identified from the research team’s extensive networks and contacts. Additional contacts made during meetings at conferences, as well as from internet searches upon suggestions from the RAG. They provided a range of services for people with ID including supported living, residential care and outreach support, and varied in size and reach, with the smallest organisations supporting under 50 people with ID within one local area, and the largest supporting over 3000 people across the UK. The most senior organisational manager was identified and contacted by email. In total, 25 services (42%) agreed to take part via written consent from a top level manager, who then provided us with the contact details of their middle managers. Whilst these 25 service represented a large sample, the phase 1 recruitment method introduced selection bias, with organisations known to the researchers more likely to be invited, and organisations where the senior manager had a positive attitude towards ‘talking about dying’ probably more likely to take part.

*Phase II: Recruiting middle managers*

The 25 senior service managers provided the contacts of a total of 728 middle managers. We contacted all of these with the invitation to send the survey link to their staff; a total of 214 middle managers responded, and between them reported that they had sent the survey link to 1130 staff members. Phase II presented us with a number of challenges, including the strong possibility of response bias, with managers probably more likely to select staff who had a particular experience of death or bereavement. It also affected our ability to get an accurate picture of the response rate, as we relied on these middle managers to let us know how many surveys they decided to send out.

This phase therefore required the most time, as it was crucial to the success of the study that middle managers sent the link to the right staff and encouraged these staff to participate. At their request, preview link was created especially for managers, allowing them to look at the survey in its entirety without answering any survey questions. This helped to encourage them to promote the survey to their staff. With regards to the contact method, we had planned to make initial contact by telephone, to be followed up by emails. However, it soon became apparent that emails were the preferred mode of communication. Many service managers were out in the field and hard to reach over the telephone; on occasions where contact was successfully made over the phone, managers were often engaged with other priorities and asked to be sent more detailed information by email. Emails therefore became the standard initial mode of contact. A follow-up telephone call was offered where managers preferred, but of the 214 managers who responded to our Phase II recruitment, fewer than five asked for a telephone call.

*Phase III: Recruiting survey respondents (direct care staff)*

A total of 724 completed the survey over a 6-month period; after excluding those who did not answer any questions beyond the first few, 690 useful responses remained. If the number of 1130 care staff who were reportedly sent the TADS link is correct, this represents a 61% response rate. On average, participants spent about 10 minutes completing the survey online; a further 20-30 minutes was spent by participants with the experience of caring for someone who had died, was terminally ill or had been bereaved. Results show that staff were willing to engage and had many opinions to share. Having managers who were supportive of the study probably helped with this.

Beyond motivating Phase I and Phase II managers, efforts to engage frontline staff were perhaps the most challenging yet most crucial, as they were ultimately our survey respondents. Since our recruitment at this stage relied on indirect contact, it was important to send a clear and engaging email that middle managers could forward to their staff at the click of a button. We believe this, in combination with a well-presented survey, was effective in engaging and motivating staff to respond.

***Elements of successful engagement of ID services and staff in research***

We found a number of interlinked strategies to be helpful for engaging ID staff at different stages of our study [see Figure 2]. The following will describe these in further detail.

*Involve stakeholders at all stages of the research process*

There was a significant networking advantage to having the RAG members involved throughout the project. This helped not only with identifying and recruiting organisations, but also to develop a realistic timeline and help the research team understand how potential participants could be approached in different types of ID services. The RAG also helped to ensure that the survey questions were of high relevance to service providers, participants, and those who would be interested in our findings. The importance of high-level organisational engagement was demonstrated by the fact that the two services from our RAG eventually accounted for about half of all surveys sent out (based on self-reports from middle managers – the research team was unable to trace the anonymous responses back to particular organisations). This was due partly to their large geographic spread and scale of their services, but also, undoubtedly, to management support for the project. The RAG members became advocates for the study within their organisations. The drawback of such support and engagement of specific services is the real possibility of response bias, as we discussed earlier.

Stakeholder engagement within the RAG was fostered by the involvement of people with ID, who required us to be creative in the way meetings were run, and needed everyone to be clear in their communication and clear about their role within the project.

At the data analysis stage, the research team presented the RAG with several possible directions for data analysis of over 200 questions from our survey. The RAG also helped to decide what kind of dissemination materials and methods would be useful to their own stakeholder groups, resulting in a range of different outputs. The RAG members were involved in developing and disseminating these materials.

Stakeholder engagement went beyond the collaboration with the RAG. Building a relationship between the research team and the participating organisations meant that there was ongoing organisational commitment to the study. This needs to be a reciprocal relationship, with the participants benefiting from the study, even if it is by understanding that their contributions will help others. The ‘giving back’ by the research team can take time, as the period between recruitment, data collection, data analysis and dissemination can be lengthy, especially in a phased study. Some of our phase I contacts were made over a year before the final analysis. The researchers remained in occasional email contact with the phase II managers, until they were able to send the final report.

*Enlist management support at all levels*

The support and engagement of managers at phases I and II was crucial. At the most senior level, the most successful recruitment was in organisations where face to face contact was made with a manager who was able to discuss and promote the study within the organisation. Some took a long time over this, waiting for the right meetings and including a piece about the study in their nation-wide internal newsletter (which we had written for them at their request). It paid off, as more responses came in after the middle managers in these organisations started sending out survey links to their staff. Key to management engagement was ensuring that all those involved felt that participating in the TADS was relevant and worthwhile to their organisation and beneficial to their staff. In phase II, much time was spent communicating with middle managers to ensure that they felt that each staff member’s participation really mattered.

We found that emails to middle managers were more likely to get a response if they were personalised, for example, by using the managers’ names several times throughout. We also included a pre-set response email which could be quickly completed with details such as the number of staff they had forwarded the survey link to; managers reported that they liked the fact that a return email was already structured for them. Additionally, providing a short and succinct follow-up email after two weeks, with a reminder of the timeframe of the study, was welcomed by middle managers who were yet to respond. Although personalising emails greatly increased the amount of researcher effort and time, it appeared to be worth it, as response rates increased. We hypothesised that the rise in responses was due to an increased sense personal responsibility and recognition of their key role in research. On the downside, this approach may also have led to over-recruitment (for example, enthusiastic managers inviting more than one staff member per home or team), but we had no way of checking this.

*Attractive and well laid-out data collection tools*

As explained above, it was crucially important to have a well-presented and attractive looking survey, making full use of technology. Free-text feedback on the survey itself was mostly positive *(“Very good survey, thank you!”; “Nice survey”)*. A few respondents reported that they found the topic upsetting *(“I found this really hard to fill out as my emotions took over and it felt as though everything I had dealt with came back and re-surfaced”)*. We received over 1,500 optional free text responses, were respondents explained their answers in further detail, sometimes at length; this suggests positive participant engagement with the survey.

*Well-structured recruitment strategy*

In developing the recruitment strategy, it is important to consider carefully who the key gatekeepers are, and who will be able to be a champion of the study within the organisation. In our study, having a three-phase recruitment strategy was time-consuming, but it was time well spent. Our strategy required significant effort and planning, with implications for budgeting and the use of researcher time.

*Time and flexibility*

It is important to note that we allocated six months for recruitment and data collection, allowing enough time for information to pass on from senior management to frontline staff members. Whilst the three-phased, top-down design of our recruitment had obvious benefits and ensured participation of the entire service, recruitment efforts were in effect tripled due to the involvement of higher managers, middle managers, as well as support workers. A six month period not only provides flexibility for middle managers to respond, but also demonstrates respect to the varying schedules and demands of both managers and their staff, accounting for annual leaves, competing work priorities; and allows time for further clarifications when needed.

Flexibility was also important. We adjusted our strategy throughout, in response to manager and participant feedback. Giving participants the flexibility to complete the survey in their own time, with the help of technology, was also useful.

*Targeted and varied dissemination*

Dissemination activities were developed together with the RAG members, who helped the research team to understand what kind of outputs or activities would be useful to their stakeholder groups. The data were analysed with support and input from the RAG. There was a vast amount of data, and we wanted to be sure we focused on the aspects that would be most useful to stakeholders. For example, with regards to survey responses to questions around “Talking about dying with people who are facing bereavement”, our advisors with ID wanted to know more about the extent to which staff talked to people with ID about this; one advisor commented, “No-one sat down with me to talk about stuff; I had an idea she [relative] was dying, but not properly.” Service managers wanted to know more about staff support: “It will be good to know what staff felt would help them”. Subsequently, in dissemination activities, these were some of the elements we focused on. A free-of-charge one day feedback conference was held in London, where staff working for ID service providers who had participated in the study were given priority to attend. All 728 managers from Phase II were sent a summary of the survey findings, along with quotes from respondents and top tips (Tuffrey-Wijne et al., 2020). In addition, family carers, people with ID and a senior manager from the RAG helped to produce short videos related to the TADS. A twitter masterclass was held over five days, followed by around 8,000 people. For example, as a result of the finding that most people with ID attend the funeral of a loved one, but very few are actively involved in it, one of the twitter threads was around involving people with ID in funerals. Twitter comments included “Thank you – I’ve never even considered this”, and “We’ve seen it so many times ‘it’s not in a person’s best interest’ to go along to a funeral, let alone be involved’.

As a result of the survey and its outcome, several ID service providers reported that they had initiated staff training on death, dying and bereavement. The process of engagement between researchers and stakeholders also helped with the identification of future areas for research.

**Discussion**

Involving ID staff as research participants not only helps to understand more about the experiences of people with ID whose own involvement as research participants may be difficult (proxy informants), but also the perspectives and practices of staff themselves.

Good levels of research participation are important. Although our recruitment efforts required a substantial amount of time and effort, it was beneficial as we achieved a good response rate of 61%, in line with what is considered the target rate (60%) (Fincham, 2008), and a higher than what is expected of a web survey (11% lower than non-web survey modes) (Fan and Yan, 2010).

The elements of success in recruiting a sufficiently large sample and engage the participants rested heavily on the engagement of ID services throughout the study. This was therefore not restricted to the period after the study design stage and before the closing of data collection period, but rather, an ongoing feature at every stage of the study. Ensuring reciprocity, by providing feedback from researchers to participants, is important (Phillipson *et al.*, 2012; Lewis and Porter, 2004).

Hall et al. (Hall, Durand and Mengoni, 2017) found supportive managers to be the key to successful recruitment, reporting on the difficulty for staff to engage with research when unsupported by senior management. They found that senior managers’ perception on the importance of research greatly determined how much time and resource were allocated, subsequently affecting whether or not staff would be able to participate at all in research. Many of our elements of success outlined above echo these findings.

Our study expands on the suggestions by Hall et al. (2017), highlighting in addition the importance of early stakeholder involvement and networking, the importance of a well-presented data collection tool, and a well-structured recruitment method. For researchers, the emphasis on fostering a RAG leading to a better designed survey and recruitment procedure not only increases respondent rates, but supports dissemination that is relevant to stakeholders. Ultimately, having the ability to transfer knowledge from the outcomes of a study to key interest groups increases the value of its findings and research efforts. For participants, there are obvious benefits of learning about results and translating the implications for practice. In our study, this spreads from the organisation level down to the individual choices each care worker makes. On the other side of the same coin, participants also affect change by informing research, fostering a greater sense of personal agency towards improved standards of practice. The increased engagement with research at all levels and in multiple directions promotes greater recognition of care organisations and staff, evidenced-based practice, and helps bridge the gap between theory and practice.

**Limitations**

All research methods and approaches to participant recruitment have their limitations, which must be considered in relation to the study’s aims and objectives. There were a number of limitations within our study design and recruitment procedures. Our initial research aims included an investigation of the proportion of people with ID within ID residential and supported living services who are diagnosed with an irreversible terminal condition, or are bereaved, within a 12 month period. Because of the anonymous nature of the study and the issue of possible double counting, we could not meet this aim. Furthermore, the reliance on gate-keepers (in our study, these were the ‘middle managers’) to select participants led to response bias and some uncertainty about precise response rates. It is important to anticipate and be transparent about such limitations, and to find ways to mitigate them if at all possible. Our participant recruitment method was suitable for investigating staff experiences and perspectives, but not for establishing prevalence of death, dying and bereavement within ID services. Future researchers who are interested in prevalence rates of particular issues or person characteristics would need to find a way that rules out double counting without compromising confidentiality.

The focus of this paper is on the extent to which our methods were effective in recruiting and engaging participants. In this, we were limited by the fact that we did not formally evaluate this. Given the importance of ‘pathways to impact’, it would be useful in future studies to incorporate a proper evaluation of the impact of various approaches to participant recruitment and dissemination strategies.

However, our analysis of what helped to improve engagement (including recruitment rates) are, we believe, transferable to other studies and other settings, both within the UK and internationally. In particular, we would advocate stakeholder involvement and ongoing active engagement between research teams and participating organisations, at all stages of research, regardless of the study methods.

**Conclusion**

Involving and engaging stakeholders throughout a research project is an important aspect of the research impact pathway. For research in ID services, these stakeholders include people with ID, family cares, service managers and frontline ID staff. Their involvement helps to ensure that the right questions are asked; data collection tools are well presented; recruitment procedures are adequate and well-structured; and, as a consequence, recruitment rates are maximised. It also helps to target dissemination activities, thus ensuring maximum research impact. Time and flexibility is required, which has resource funding implications.

In our study, ID staff were invaluable and enthusiastic advisors and participants, who were able to provide unique insight into best practice for talking about loss with people with ID. Bridging the research-practice gap in the field of ID will ultimately lead to improved support for people with ID. Frontline ID staff can play a crucial role in the bridging of this gap.

**References**

Bekkema, N., De Veer, A., Wagemans, A., Hertogh, C. and Francke, A *.* (2015) ‘“To move or not to move”: A national survey among professionals on beliefs and considerations about the place of end-of-life care for people with intellectual disabilities’, *Journal of Intellectual Disability Research*, 59(3), pp. 226–237. doi: 10.1111/jir.12130.

Fan, W. and Yan, Z. (2010) ‘Factors affecting response rates of the web survey: a systematic review’, *Computers in Human Behavior*, 26(2), pp. 132–139. doi: 10.1016/j.chb.2009.10.015.

Fincham, J. (2008) ‘Response rates and responsiveness for surveys, standards, and the Journal’, *American Journal of Pharmaceutical Education*, 72(2), p. Article 43. doi: 10.5688/aj720243

Forrester-Jones, R. (2013) ‘The road barely taken: Funerals, and people with intellectual disabilities’, *Journal of Applied Research in Intellectual Disabilities*, 26(3), pp. 243–256. doi: 10.1111/jar.12022.

Hall, N., Durand, M. and Mengoni, S. (2017) ‘“...their opinions mean something”: Care staff’s attitudes to health research involving people with intellectual disabilities’, *British Journal of Learning Disabilities*, (May), pp. 1–10. doi: 10.1111/bld.12195.

Hoover, J., Markell, M. and Wagner, P. (2005) ‘Death and grief as experienced by adults with developmental disabilities: initial explorations’, *OMEGA: The Journal of Death and Dying*, 50(3), pp. 181–196. doi: 10.2190/KAHM-0RL8-NF59-8LRN.

Hunt, K., Bernal, J., Worth, R., Shearn, J., Jarvis, P., Jones, E., Lowe, K., Madden, P., Barr, O., Forrester-Jones, R., Kroll, T., McCarron, M., Read, S. and Todd, S*.* (2019) ‘End of life care for people with intellectual disability: a retrospective cross sectional UK study’, *BMJ Supportive and Palliative Care*, Epub ahead, pp. 1–9. doi: 10.1136/bmjspcare-2019-001985.

Lewis, A. and Porter, J. (2004) ‘Interviewing children and young people with learning disabilities: guidelines for researchers and multi-professional practice’, *British Journal Of Learning Disabilities*, 32, pp. 191–197. doi: 10.1111/j.1468-3156.2004.00313.x

Lord, A. J., Field, S. and Smith, I. C. (2017) ‘The experiences of staff who support people with intellectual disability on issues about death, dying and bereavement: A metasynthesis’, *Journal of Applied Research in Intellectual Disabilities*, 30(6), pp. 1007–1021. doi: 10.1111/jar.12376.

Northway, R., Todd, S., Hunt, K., Hopes, P., Morgan, R., Shearn, J., Worth, R. and Bernal, J*.* (2018) ‘Nursing care at end of life: a UK-based survey of the deaths of people living in care settings for people with intellectual disability’, *Journal of Research in Nursing*, online fir, pp. 1–17. doi: 10.1177/1744987118780919.

Phillipson, J., Lowe, P., Proctor, A. and Ruto, E*.* (2012) ‘Stakeholder engagement and knowledge exchange in environmental research’, *Journal of Environmental Management*. Elsevier Ltd, 95(1), pp. 56–65. doi: 10.1016/j.jenvman.2011.10.005.

Read, M. (2018) *The research impact handbook*. 2nd edn. Huntley: Fast Track Impact.

Ryan, K., Guerin, S., Dodd, P. and McEvoy, J*.* (2011) ‘Communication contexts about illness, death and dying for people with intellectual disabilities and life-limiting illness’, *Palliative and Supportive Care*, 9(2), pp. 201–208. doi: doi.org/10.1017/S1478951511000137.

Stancliffe, R. J. *et al.* (2016) ‘Knowing, planning for and fearing death: Do adults with intellectual disability and disability staff differ?’, *Research in Developmental Disabilities*. Elsevier Ltd., 49–50, pp. 47–59. doi: 10.1016/j.ridd.2015.11.016.

Todd, S. (2013) ‘“Being there”: the experiences of staff in dealing with matters of dying and death in services for people with intellectual disabilities.’, *Journal of Applied Research in Intellectual Disabilities*, 26(3), pp. 215–30. doi: 10.1111/jar.12024.

Tuffey-Wijne, I. and Rose, T. (2017) ‘Investigating the factors that affect the communication of death-related bad news to people with intellectual disabilities by staff in residential and supported living services: An interview study’, *Journal of Intellectual Disability Research*, 61(8), pp. 727–736. doi: 10.1111/jir.12375.

Tuffrey-Wijne, I. (2013) *How to break bad news to people with intellectual disabilities: a guide for carers and professionals*. London: Jessica Kingsley Publishers.

Tuffrey-Wijne, I., Bernal, J. and Hollins, S. (2008) ‘Doing research on people with learning disabilities, cancer and dying: ethics, possibilities and pitfalls’, *British Journal of Learning Disabilities*, 36(3), pp. 185–190. doi: 10.1111/j.1468-3156.2008.00519.x.

Tuffrey-Wijne, I., Finlayson, J., Bernal, J., Taggart, L., Lam, C., Todd, S. (2020) Communicating about death and dying with adults with intellectual disabilities (ID) who are terminally ill or bereaved: a UK-wide survey of ID support staff. *Journal of Applied Research in Intellectual Disabilities* online first, doi: 10.1111/jar.12714

Wiese, M., Dew, A., Stancliffe, R., Howarth, G. and Balandin, S*.* (2013) ‘“If and when?”: the beliefs and experiences of community living staff in supporting older people with intellectual disability to know about dying’, *Journal of Intellectual Disability Research*, 57(10), pp. 980–992. doi: 10.1111/j.1365-2788.2012.01593.x.

Figure 1: ***Recruitment to the “Talking About Dying Survey”***

Top level management identified and sent Phase I Participant Information Sheet and short survey form to complete and return (n=25; 42.4%)

ID service providers identified and contacted by email or phone (n=59; 100%)

Did not reply or call back (n=26)

Asked for more information but did not provide next person of contact (n=8)

Middle managers contacted by email or phone and emailed Phase II Information Sheet; managers asked to choose at random 1 care worker per team they oversee (n=728; 100%)

Did not reply email detailing how many care workers were forwarded the email (n=508)

Middle managers deemed survey not applicable to service (n=6)

Middle managers (n=214; 29.4%) forwarded Phase III email with survey link and Participant Information Sheet to care workers (n=1130; 100%)

Total surveys completed (n=724; 64.1%)

One participant was excluded as she did not meet inclusion criteria. She cared for a child under the age of 16 and therefore could not be categorized as an adult. (n=1)

Participants who did not answer beyond Question 5 of the survey were excluded. (n=33)

Total surveys analysed (n=690; 61.1%)

 Figure 2: ***Elements of successful engagement of ID services and staff in research***