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The development of a mentor training program for practitioners working with autistic adults

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Abstract

This study focuses on the development and evaluation of a training program for would-be mentors of adults on the autism spectrum. Relevant literature in this field is reviewed along with existing training and mentoring services for autistic adults, and the development of a mentor training programme described. The training programme was developed by the project team with advice from the project advisory and steering groups. This thesis reports on the development of participant feedback as to the effectiveness of the training model devised. Unlike many contemporary mentoring models in the field of autism, this project took a social model approach with training informed, designed and delivered with significant input from autistic people. In participant feedback, this aspect of the training was the best received. Following feedback from participants however, it is recommended that this training model be developed to be covered over a two-day period and contain added emphasis on the reliability and consistency of mentors, role boundary setting, and the facilitating of goal-setting.
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Dedicated to the continuation of the Participatory Autism Research Collective (PARC)
Section 1: Introduction

1.1 A note on terminology

There is much current debate regarding the terminology related to autism. This article will resist people first phrasing, in accordance with other autistic voices (Sinclair, 1993; Sainsbury, 2000):

“We are not people who “just happen to have autism”; it is not an appendage that can be separated from who we are as people, nor is it something shameful that has to be reduced to a sub- clause.” (Sainsbury, 2000: 12).

1.2 Project rationale

Until recent decades autism was considered to be a rare developmental ‘disorder’ that affected a small proportion of the population. However, with the widening of diagnostic criteria, the numbers of those now considered to be somewhere on the autism spectrum is estimated to be around 1 in 100 (NAS, 2016a). Although prevalence figures for autism are hard to judge, due to the changeable criteria and diagnostic practices that it has undergone historically, and due to autism being diagnosed using behavioural characteristics, an accurate figure is not possible, as there is much contention over what exactly autism is (Milton, 2012a; Runswick-Cole et al. 2016). What can be said however, is that at least the majority of those now considered to be on the autism spectrum are those without significant learning disabilities and who are often highly articulate in their communications. However, research in the field of autism is primarily concentrated on a medical model framework and concentrated on children and young people (Pellicano et al. 2015).
Research into, as well as practical support, for the needs of adults on the autism spectrum is sparse, and the nuanced needs many have within educational and work settings can lack recognition. At a forum held by Research Autism in 2007 entitled ‘Successful Futures for Adults with Autism’ participants highlighted difficulties experienced with navigating social life, including: managing their own practical and financial affairs, accessing education and training opportunities, securing and maintaining employment, and maintaining good physical and mental health. There was a common feeling expressed that existing models of support for adults on the autism spectrum, which often involve being part of a large group of people, were not helpful. Many described how they felt stressed or unsure in such surroundings, preferring a one-to-one relationship which could then be broadened over time. Many said that they would only want this support on a time-limited basis, but that it should be goal-oriented, specialised and based on a personal life coach or mentor model. Many participants said they would like to use the allowances they received for personal support to pay for such services, but few had access to such services in their locality.

Due to these findings, Research Autism later acquired funding to undertake a two-year pilot study to establish a mentoring scheme, designed with significant input from autistic people and their families, in order to evaluate the programmes effects on the well-being of the participants and progression toward their stated goals. The aims of this study were to two-fold and included subordinate aims as follows:

Aim one: to develop a sustainable research-informed mentor training programme:

1a: To review existing training for mentors of people on the autism spectrum.

1b: To develop a mentor training programme with input from adults on the spectrum.
Aim two: to assess the effectiveness of the mentoring programme developed for adults on the autism spectrum:

2a: To assess whether mentoring impacts on goal achievement and satisfaction for mentors and mentees.

2b: To assess whether it has any impact on the quality of life of both mentors and mentees.

2c: To use qualitative interviews to gain an understanding of participants’ perceptions of the mentoring programme in order to improve the programme based on the views of those involved.

This dissertation report concentrates on the first of these project aims, of reviewing existing training and mentoring services for autistic adults, and the development of a mentor training programme. The remit of the project included the production of a one-day training course for would-be mentors of autistic adults. This training programme was developed by the project team with advice from the project advisory and steering groups. This thesis reports on this initial phase and concentrated on the following research questions:

What information and activities are most effective for meeting the training needs of would-be mentors of autistic adults?

What aspects of a training programme for would-be mentors of autistic adults are most appreciated by those undertaking the training?

This dissertation report is structured into six sections. Following this introduction is a literature review which considers the mentoring role, relevant literature on autism and how this related to the mentoring of autistic people and literature regarding the use of Personal
Construct Theory in the design of the mentoring program and accompanying training materials. In section 3, the methodology of this dissertation is discussed, followed by section 4 regarding the research methods employed. Section 5 gives an overview of the results, whilst section 6 discusses and concludes the main findings and limitations of the study, before giving recommendations for continued research in the area.
Section 2: Literature review

The aspect of the mentoring project being focused on in this dissertation report set out to answer the following questions:

What information and activities are most effective for meeting the training needs of would-be mentors of autistic adults?

What aspects of a training programme for would-be mentors of autistic adults are most appreciated by those undertaking the training?

In this section, a review of relevant literature is given regarding mentoring, autism, research in the area of mentoring autistic people, and lastly in the area of Personal Construct Theory (PCT) which through the literature review became influential on the mentoring model and training adopted within this project.

2.1 Literature regarding the mentoring role

According to The Mentoring and Befriending Foundation (2014), mentoring is a time-limited goal-orientated relationship that supports both personal and vocational learning and development. It involves an experienced person providing guidance and support to another (less experienced) person through a variety of methods (Western 2012). Western (2012) suggest that it is a necessity for a mentor to have an understanding of the social world and perspective of their mentee, as well as the skills to share their experience in an effective and helpful way.

Mentoring can occur either formally or informally (Miller 2002). Informal or ‘natural’ mentoring can be said to develop within all kinds of social relationships, whereas formal or
planned mentoring operates within a structured programme with clear objectives, where mentors and mentees are matched. Planned mentoring can vary vastly depending on the institution in which it is delivered and the philosophy guiding its practice, but there are usually some shared characteristics, described by Miller (2002) as including:

- It is a deliberate, conscious and voluntary relationship.
- It may or may not be time limited.
- It is supported by an organisation.
- It occurs between an experienced person and one or more other (less experienced) persons.
- It is non-hierarchical.
- The relationship is expected to be beneficial to both mentor and mentee.
- It will typically include elements of interpersonal support, guidance, mutual exchange, sharing of wisdom, and coaching.

Despite these characteristics, both mentors and mentees are involved in negotiating the form of their responsibilities within the relationship and are therefore involved in defining what mentoring is and what mentors do (Miller 2002).

Within the initial meeting between mentor and mentee the initial parameters of the relationship are established. The mentor and mentee need some time to start to plan what the goals of the mentoring experience are to be. Commitment to the mentoring relationship from both parties is essential for its potential success, but not always an easy process to sustain. According to The Mentoring and Befriending Foundation (2014), distinctions need to be made between mentoring and other support roles such as coaching. Whilst there are
similarities between mentoring and coaching and other support roles such as befriending, advocacy and counselling, there are also important differences that need to be addressed when setting the parameters of the mentoring relationship. As an example, in both mentoring and coaching roles, one may work toward set goals and objectives, with outcomes potentially being measured in some way with regard to improved performance of a task. However, a mentoring relationship is often more open-ended, with the mentor more in the role of facilitating the learning of their mentees, as well as more holistic goals such as self-discovery or personal growth. Within mentoring relationships it is vital to maintain professional boundaries for the sake of both parties. The outcomes of a mentoring relationship are often broadly defined and reflect the mentees progress in their development toward a negotiated goal. Such a partnership can be time-limited or long-term, and can evolve into a mutually beneficial collaboration where mentors can learn as well from the feedback, insights and reflections of their mentees. The positive outcomes of a mentoring relationship can be seen to be largely dependent on the quality of the mentoring experience and the interactions between mentor and mentee.

2.2 Literature regarding autism

In order to develop a mentoring programme for autistic adults, it is necessary to look at what aspects of autism people need to know about in developing their mentoring expertise. Whilst a person-centred ethos is essential for a mentoring programme, there may be many aspects of the autistic experience which could impact on mentoring relationships, and could act as a foundational knowledge to reflect upon for would-be mentors working with autistic adults.
The term autism was first used by a psychiatrist called Bleuler in 1911 to try and describe a type of what was then called ‘childhood schizophrenia’. His descriptions however, only show a passing resemblance to how autism is thought of today:

"The schizophrenics who have no more contact with the outside world live in a world of their own. They have encased themselves with their desires and wishes...they have cut themselves off as much as possible from any contact with the external world. This detachment from reality with the relative and absolute predominance of the inner life, we term autism." (Bleuler, 1911, cited in Parnas et al., 2002: 131).

In the 1940’s two psychiatrists called Kanner, and working separately Asperger, were like Bleuler, studying small groups of children deemed as having some form of ‘childhood schizophrenia’. Both found that with the groups of children they were working with a set of distinct symptoms were being identified that were markedly different from schizophrenia as it was conceived of at the time. Kanner’s work laid the foundation for early accounts of autism, whilst the work of Asperger was left largely undiscovered until the 1970s. Definitions of what autism is, and also what caused an autistic developmental pattern in children have been hotly contested ever since, including an unfortunate era where autism was thought to be a reaction to ‘refrigerator mothers’, a theory long since shown to be totally untrue.

One of the most important developments in the history of autism in Britain was the work of Wing and Gould (1979) and the subsequent widening of the autism spectrum to include Asperger syndrome. This work largely created the contemporary definition of autism as a ‘triad of impairments’ in: social communication, social interaction, and imagination
(repetitive interests/activities). Since this time, diagnostic systems have changed to reflect these changes in definition.

The most commonly utilised definition of autism that one sees today is that it is a:

“...lifelong developmental disability that affects how a person communicates with, and relates to, other people. It also affects how they make sense of the world around them.”

(NAS, 2016a).

Although this definition of autism is much contested, such a definition of behavioural deficit and impairment has come to characterise both the DSM-V (2013) and ICD-10 (1992) diagnostic criteria. Autism is thus diagnosed according to ‘qualitative’ impairments in all three areas.

In terms of theorising the ways in which autistic people make sense of the world around them, explanations have been dominated by cognitive theories framing autistic ways of thinking in terms of deficit and impairment. One of the most commonly cited theories suggests that autistic people can have a deficit in executive functioning. This phrase refers to the cognitive ability to maintain an appropriate problem-solving strategy needed in order to complete a task. Although first-hand accounts of autistic people often reference difficulties in switching attention, or adverse reactions to interference with attention (e.g. Tammet, 2006), such a theorising of autistic ways of thinking are not without their critics. Some on the autistic spectrum can perform well in executive functioning tasks, whilst people with other medically defined conditions may also struggle with such tasks, suggesting a lack of specificity to autism. It may be the case that such differences are better
explained by theories such as a weak drive toward central coherence and monotropism theory.

Happe (1994a) suggested that another difficulty with executive functioning theory is the lack of explanatory value regarding the strengths and talents that are often displayed by autistic people. Shah and Frith (1983) found that autistic people outperformed non-autistic people in tasks such as the embedded figures test, or picking out details from a visual array. Shah and Frith (1983, 1993) suggested that autistic people are often better at tasks that require the ignoring of ‘overall meaning’ in order to complete. Happe (1994a) suggested that autistic people thus had a ‘weak drive toward central coherence’ in terms of perceptual meanings grasped from any given context, whilst simultaneously having the ability to process fine details within their perception. A difficulty with this theory is that autistic people are able to process contextual meanings, yet it may be the case that in some contexts there may be a cognitive tendency to focus on particular aspects of a situation, rather than others.

Murray et al. (1992; 2005) argue that rather than framing autistic cognition in terms of a dysfunctional deficit, the central core feature of such ways of thinking would be the atypical strategies employed by autistic people in their distribution of attention, or perceptual interest. This theory has been called ‘monotropism’ or an ‘interest’ model of autism. This theory would suggest that the amount of attention available to any individual at one time is limited, and that the shaping of cognitive processes depends on a competition between mental processes for this scarce resource. Murray et al. (2005) argue that the way in which attention is utilised by people is normally distributed from those who concentrate deeply on a small number of interests and those with a diffuse but wide attention strategy, akin to a
‘torch beam’ or a ‘diffused light’, with those at the tightly focused end of this spectrum can be said to be autistic. Murray et al. (2005) suggest that social interaction, the use of language, and the shifting of object attention (implicated by other psychological theories) are all tasks that require a broad attention, and are inhibited by a narrow use of attention.

This theory highlights aspects of the autistic experience not explained sufficiently by the dominant psychological theories in the field. For instance, how individuals on the autism spectrum show a tendency toward either being passionately interested in a task or phenomena, or not interested at all, or how an unanticipated change can lead to a catastrophic disconnection from a previously safe state of mind. When employing a monotropic interest perceptual strategy, the ability to use previously learnt information and schema may be compromised, as information can be channelled along a limited range of interests.

“We suggest that the uneven skills profile in autism depends on which interests have been fired into monotropic superdrive and which have been left unstimulated by any felt experience.” (Murray et al. 2005: 143).

An often commented upon aspect of the autistic experience is that of having a ‘spiky’ or ‘uneven’ profile of abilities and interests. This aspect of autistic experience is understandable through the theoretical perspective of monotropism, yet is often unrecognised by service providers and support workers, particularly if they are reliant on categorical notions of the cognitive functioning of autistic people. Highly verbal or articulate autistic people may be perceived to be capable in areas in which they struggle, and those with less verbal ability may be assumed to be lacking in skills and learning capacities.
Following the theory of monotropism, social interaction with other people may be somewhat dependent on connections being made in the fulfilling of passions and interests that the autistic person shows. From such a perceptual viewpoint, difficulties understanding the perceptions of others may become apparent, but it can also be said that non-autistic theory of autistic mind can leave something to be desired (Milton, 2012a).

Perhaps the most dominant theory in the study of autism is that autistic people have a deficit in, or even lack a ‘theory of mind’ (Baron-Cohen et al. 1985). The concept of theory of mind refers to the ability to empathise with other people and imagine their thoughts and feelings, which can then be used to comprehend and predict their likely behaviour. Baron-Cohen et al. (1985) initially based this theory on the finding that 80% of their sample of autistic children between the ages of six and sixteen failed at false belief tasks such as the ‘Sally-Ann’ test. These findings were repeated in subsequent studies that utilised real people instead of toy dolls (Leslie and Frith, 1988).

This theory has been criticised on a number of levels however: Task failure on false-belief tasks could be due to difficulties in language processing or memory (Eisenmajer and Prior, 1991), or a lack of motivation to deceive (DeGelder, 1987). Some theorists questioned its applicability to all on the spectrum when 20% of children passed such tests, or its value for explaining all the aspects of what constituted the autistic difference (Happe, 1994a). An impaired theory of mind deficit as a general theory of autism has since been revised in order to differentiate between an ability to ascertain the feelings of others, and the development of affective empathy once those feelings are recognised. Later studies by Baron-Cohen (1992) and Happe (1994b) also found that the ability to successfully complete theory of mind tasks increased with age and IQ, suggesting a delayed ‘mentalising’ capacity.
Autistic scholars, such as Milton (2012a; 2014) and Chown (2014) have argued that a so-called deficit in social functioning cannot solely be located within the mind of an individual, and that in social interactions between autistic and non-autistic people a breakdown in understanding can easily occur in both directions due to the differing dispositions and perceptions of the parties involved. As coined by Milton (2012a), a ‘double empathy problem’ exists between the two interaction partners, as both can have significant challenges in understanding the intentions of each other.

An extremely important aspect of the autistic experience is that of sensory sensitivities. This feature of autism was only recently recognised within diagnostic criteria (DSM-V, 2013). Autistic people often report both hyper and hypo sensitivities to all sensory sensations. Such sensitivities can vary from person to person and across contexts. With a fragmented or monotropic cognitive style, sensory integration for autistic people can also be a challenge, with a high level of crossover with those diagnosed with Dyspraxia.

Due to the sensory and social experiences that autistic people experience, many also report high levels of stress. Experiences of stress are certainly not unique to autistic people, yet problems associated with chronic stress levels are common. Without available options or the ability to communicate when highly stressed, reactions can lead to what can be characterised as ‘meltdowns’ (emotional outbursts), ‘shutdowns’ (characterised by withdrawal), and panic attacks. Such reactions can be seen as fight, flight or freeze type of responses, yet may be misinterpreted by others due to issues such as the double empathy problem (Milton, 2012a).

The dominant theories of autism, much like the diagnostic criteria for autism, locate the challenges faced by autistic people primarily within the brains/minds of the autistic person,
rather than in the social milieu within which they might find themselves, or in the relationships they have with others. Milton (2012a) argues in contrast however, that the social subtext of a situation is never a given, but actively constructed in the interactions people have with one another.

Within wider disability activism and critical disability studies, there are a number of models that have been explored in regard to the study of disability (the two most frequently referenced being the traditional medical and social models). The traditional medical model of disability would view disability as a set of problems or deficits belonging to the individual and in need of remedial intervention. This view has been widely criticised by disability activists (Oliver, 1990). In contrast to the medical model is the social model of disability which sought to socially situate the experience of disability. The traditional social model of disability (e.g. Oliver, 1990) demarcates socially situated disability from biological impairment. However, the notion of impairment also being socially constructed has been explored by disability scholars (Milton, 2015; Sherry, 2016). Within the field of autism, it can be seen that medical and cognitive models dominate the discourse. Yet, this pattern is reversed within the autistic community, with many autistic adults expressing a view more akin to a social model approach (Milton, 2016a).

2.3 Relevant research regarding mentoring for autistic people

Both Access to work and student mentoring for people on the autism spectrum is available through various schemes in the UK, yet specialist schemes are rare and research on the topic rarer still. The only area of mentoring for people on the autism spectrum to have begun to gain the attention of researchers has been student mentoring schemes for College and University students. Gelbar et al. (2014) conducted a systematic review of articles
describing the experience and support schemes made available for people on the autism spectrum attending College or University. This review only found twenty articles referring to sixty-nine people in total, with only two of these studies being ‘experimental’ in nature, neither of which were evaluating a mentoring scheme. The other eighteen studies were all individual case study reports. Such a scarcity of research into the area indicated a genuine need for further study.

Gelbar et al. (2014) found that the majority of the studies looking into experiences of post-compulsory education included accounts of isolation and loneliness, and problematic mental health. Much guidance that has been written for preparing and supporting people on the autism spectrum in post-compulsory education has been done so from a clinicians perspective, suggesting interventions to address academic modifications, independent living and social skills, vocational goals, and mental health supports in order to improve the quality of life of such students (Van Bergeijk et al. 2008). There are some potentially valuable insights in such accounts, such as smaller settings and class sizes, and utilising strengths and areas of interest. By taking a medicalised deficit model view of autism, however, they also recommend strategies such as role playing, generalising, teaching people on the spectrum to recognise their own emotions through explicit instruction, and so on, which would be much criticised by a number of autistic scholars and activists (Milton, 2012a, 2014).

Mentoring can be seen as the development of a one-to-one relationship that helps an individual to learn from their own experiences as well as from the experiences of another, with both mentors and mentees potentially gaining from the experience. In the context of
autistic people however, the benefits from such a relationship may be difficult to attain due to the double empathy problem (Milton, 2012a; 2014; Chown, 2014).

Brown et al. (2010) examined evidence regarding mentoring practices for students with disabilities in post-compulsory education. They found that literature in the area was minimal, with only ten pieces of research meeting their criteria for review. A number of themes did emerge from this review however, including: the potentially useful role of technology, focusing on particular disability groups, the usefulness of mentoring for academic, career and social ‘skills’ guidance, and the value of establishing a long-term mentoring relationship. Brown et al. (2010) conclude by stating that mentoring relationships with students with disabilities have the best opportunity for success where there is a flexible and multi-layered system of support in place.

There is some contention within the literature however regarding the goals of mentoring and who should set them. Whilst the Mentoring and Befriending Foundation (2014) suggest that goals should ideally be set by mentees, Griffith et al. (2012) argue that little is known about what extent people on the autism spectrum need supports to help ameliorate the ‘core characteristics of the condition’. Such a framing places autism within a deficit model in need of remediation from the outset. Griffith et al. (2012) suggest that presenting a ‘false self’ to others, despite being difficult, could be a useful strategy, and for some was easier than ‘being themselves’. It needs to be said however, that masking one’s identity can have significant negative effects on autistic well-being. Such negative consequences include the possibility of falling to ‘pass’ anyway, succeeding with great effort, but this potentially leading to inaccurate expectations being placed on people by others, the potential for exhaustion, and so on. Griffith et al. (2012) present a model of support based on a
behavioural deficit and remedial model of autism and could be argued to mis-frame the issues faced by their participants, or at least they ignore alternative explanations. For example, they suggest that participants were concerned about how they might be perceived to be ‘odd’ by other people, which in turn could exacerbate their feelings of anxiety and depression, whilst others ‘implied’ that autism was an affliction placed upon them, separate to their ‘core’ or ‘real’ sense of self. Instead of seeing these examples as evidence of potential internalised stigma and psycho-emotional disablement (Reeve, 2011; Milton and Moon, 2012), the authors suggest that ‘passing’ and ‘masking’ skills are of social benefit to autistic people and may reduce proneness to anxiety and depression. In the view of Milton and Moon (2012), such a view would be clear example of normalisation and cognitive ableism and could lead to exacerbating the very problem such a behavioural model is attempting to remediate. Whilst Griffith et al. (2012) acknowledge that other participants saw being autistic as freeing them from the constraints of ‘normal thinking’; they seem to have had more of a difficulty themselves in this regard. Griffith et al. (2012) argue that autistic people and their families should be actively consulted and included and have a valued role in relevant contexts. From the point of view of community-based participatory research however, such efforts at inclusion fall a long way short of the ideal.

Curtin et al. (2015) conducted a small pilot study regarding the feasibility and efficacy of an individualised mentoring program for teenagers on the autism spectrum. In this program, mentees met with mentors for two hours a week over a six month period. Like Griffith et al. (2012), they introduce a normative approach through the program described, with a focus on the teaching of social skills and competencies, framed by Curtin et al. (2015) as critical for independence in adulthood. The learning of normative social skills were assumed by Curtin
et al. (2015) to lead to better quality of life and well-being. Such work ignores the criticisms that autistic people make of normative social skills training and academic work that critiques such a normative approach (Milton, 2016a). Curtin et al. (2015) did have goals set and how to reach them through mutual discussion between mentor and mentee, and a clear distinction was made on the program between mentoring and befriending services. The goals set were constrained however to five core areas: self-esteem, healthy relationships, independent living, community involvement, and education/vocation. It is questionable as to what extent structuring goal setting to this extent is really person-centred. The mentors on this program received training in an overview of autism (‘spectrum disorders’), communication skills, behavior management, and role modeling ‘proper social behaviour’. Training in autism that ignores social model approaches and the voices and concerns of the autistic community may lead to unhelpful or even counterproductive practices being implemented. Despite these issues, Curtin et al. (2015) found a general trend toward improvement in self-esteem and quality of life measures for their participants. They also found that the mentoring worked best, when the pairings worked on the goals that mentees had primarily set for themselves. Issues were also found with regard to the boundaries of the mentoring role, with one participant asking their mentor if they were ‘dating’.

The normative trend in studies regarding mentoring and autistic people is continued by Arnes et al. (2015). This study evaluated a mentoring program designed to help students navigate both social and academic aspects of post-compulsory education. Despite presenting a highly normative model of autism, the authors attempted to locate the study within disability theory and also highlighted issues regarding a sense of belonging within a University setting, such as safety and the building of a network of peers. In this program,
matched pairings would meet either weekly or bi-weekly, with mentors drawn from graduate psychology students and supervised by a clinical psychologist.

A contrasting study to the standard normative approach was reported by Beadle et al. (2013) which suggested that autistic people themselves were in the best position to offer insights to others about why autistic people acted in the way that they may do, and that all services for autistic people should have significant input from autistic people. Such calls for inclusion and participation have also been mirrored by autistic writers within the context of research practice (e.g. Milton and Bracher, 2013). Beadle et al. (2013) report on a project that was set up to enhance the presentation skills of young autistic adults who expressed an interest in talking publicly about their lives. Whilst a specific goal and one open to criticism from within the autistic community for potentially presenting autistic people as ‘self-narrating zoo exhibits’, it is one often found amongst young autistic adults and an interest that can be exploited. Their project involved a series of mentoring sessions and drew heavily of participatory research methods to determine the content of the sessions. After each session, each participant was invited to record their thoughts in reflective journals. The project received a great deal of positive feedback from all involved in it, in terms of having a positive experience, gaining confidence, and learning new skills.

In order to try and build bridges across the divide that can occur in social interactions between autistic and non-autistic people (Milton, 2012a), tacit understanding, trust and rapport, need to be nurtured over time. Thus, personal qualities such as humility, patience and willingness to listen are paramount. Imposing interpretive frameworks and normative ideologies onto autistic people are likely to be less successful. A recent study by Hamilton et al. (2016) found that good quality training was essential to University mentors working with
autistic people. With significant input from autistic people, such training could be even more impactful and productive.

Where studies have used wellbeing indicators with adults on the autism spectrum, one often finds much lower self-assessments (Bracher, 2014). Yet, it is debatable to what extent standardised measures of wellbeing capture autistic experience and sensibilities. Current measures of wellbeing used in the context of autism have been developed with a non-autistic population and hence, may not adequately reflect an autistic perspective. Therefore, this area of research has been gaining increasing attention (Jones and Hurley, 2014). A number of studies have shown the potential of utilising Personal Construct Theory (PCT) (Kelly, 1955, Salmon, 2003) with autistic populations however for gaining a clearer insight into the perspectives of autistic participants (Moran, 2006; Williams and Hanke, 2007; Milton, 2012b; Greenstein, 2013).

2.4 Literature regarding the use of Personal Construct Theory (PCT)

PCT developed as a pragmatic theory through George Kelly’s (1955) psychotherapy practice. At the time of conception, such therapy was dominated by two divergent schools of thought: Psychoanalysis and Behaviourism. Both perspectives were vastly different to one another, yet both took the standpoint that people were moved to act by forces largely outside of their own control (the unconscious mind and environmental influences respectively). In contrast, PCT saw the person as an agent, making choices and decisions and acting upon them. This conceptualisation would not divorce actions from the context within which people act, but for Kelly (1955), it was the constructions that an individual places on events that shapes the meanings they form and the reactions they have to events.
Thus, the starting point for PCT was the idiosyncratic ways in which people make sense of the world and how that leads to social action.

Such an approach to personal constructions draws heavily on a phenomenological approach, attempting to approach issues through the viewpoint of the individual experiencing them, rather than fitting them into a priori theories. Kelly (1955) used the term ‘constructive alternativism’ to suggest that there were many differing ways of perceiving and making sense out of the same thing or event, and rather than seeing any interpretation as correct, one should look pragmatically at how useful such a framing is to one’s purposes. For Kelly (1955), following on from the work of Mead (1934), social roles were not fixed positions, but something navigated by an individual in their interactions with others. Importantly in the context of autism, it involved placing oneself in the position of the other with whom one was interacting, so that one could adjust one’s social performance accordingly:

"He [George Kelly] argued that there were two ways of treating other people. You can relate to others in the way that the early behaviourists thought normal, and treat them as 'behaving mannequins'. Only psychopaths do this, he claimed. The moral way to relate is to act in the light of the other person's view of things. In other words, taking their thoughts and feelings into consideration." (Butt, 2008: 13).

Kelly (1955) envisaged the personal construct to be ways of construing events along bipolar continuums, e.g. from happy to sad, anxious to relaxed. This is not to say constructions are of the nature of either/or extremes, but can be placed along continuums. Placed together these constructions comprise a ‘construct system’. In this sense, discourse tells us little about the actual event, but tells us a lot about how someone is construing an event.
Constructs are more than just conceptualisations however, as they are both ways of reflecting upon phenomena and of motivating social action. Construing can also be seen as something that is an active process, rather than something static that one ‘has’. Therefore, a construct system is not a cognitive entity existing in a vacuum, but is socially and discursively situated.

According to PCT, there is no such thing as a static ‘self’, as a cognitive entity made up of ‘traits’. Equally, an individual is not seen as an empty vessel moved to act by outside forces alone, rather that there was a direction to a person’s actions, or as later post-modernists may have said, a ‘line of flight’ (Deleuze and Guattari, 1980). To make sense of this phenomenon, Kelly suggested that people develop ‘core constructs’ at the centre of their construct systems, which are therefore essential ways in which people construe the ‘self’. In this conceptualisation, the ‘self’ is neither static nor fluid, not working with a psychological vacuum nor totally driven by external forces, but as a personally constructed ‘clumping’ of meaning-making activity (Milton, 2013; 2013a; 2013b; 2016b), a ‘self-theory’.

“The rules of logic do not apply in a person’s phenomenology. Instead, we see an idiosyncratic psycho-logic in operation.” (Butt, 2008: 41).

Another technique developed utilising PCT was that of ‘the Salmon line’ named after its creator Phillipa Salmon (2003). This technique asks an individual to draw a line with words representing opposite extremes at each end (e.g. anxious to relaxed). The individual is then asked to place themselves along this line and where they would like to be in future. The individual is then asked to write or talk about how they think they will get from one point to the other. It is in this discursive space that for Salmon (2003), the learning experience takes place. Similar techniques have also been devised with regard to expressing a sense of ‘real’
and ‘ideal’ self-image (of course with both being seen as constructions), or of the organisation one works within. Such techniques have helped inspire participatory work with autistic children (Moran, 2006; Williams and Hanke, 2007; Greenstein, 2013).

Williams and Hanke (2007) employed PCT methods to explore with autistic children how they experienced school life and imagining how it might be different. They argued that pupils should be involved as much as possible in making decisions about issues that most directly affected them. Williams and Hanke (2007) recognised that they needed practical tools with which to gain a genuine picture of the pupil’s views without pre-determining what these might be. This led to methods being used with the children such as building a model of their ideal school.

PCT examines how a person construes and constructs themselves in narrative meaning (and themselves in relation to others). Within this theory, people were initially regarded as like everyday ‘scientists’, where they develop theories about the world and then test them out in real life situations. Although this idea can be critiqued, as people are often not very conscious about the decisions they make and the meanings they form, as Moran (2006) points out, this way of meaning-making may be more akin to some autistic sensibilities. By focussing on how an individual construes the world, idiosyncratic constructions are anticipated, and if understood better, can help practitioners to engage with the autistic people that they work with.
Section 3: Methodology

In section 2, literature regarding mentoring, the nature of autism, and the links between autism and mentoring was all reviewed. In the following section the methodological rationale for this study is outlined, followed by a report regarding ethical considerations and a positionality statement.

3.1 Methodological rationale

It has been argued that traditional research in the social sciences has perpetuated unequal power relationships experienced by groups of people who have historically experienced marginalisation in society, with research being done ‘to’ rather than ‘with’ people (Barnes & Sheldon 2007). The ‘emancipatory’ research paradigm purports that the participants of research should have involvement with and control over the research agenda and process (Barnes & Sheldon 2007). Furthermore, the overall aim of emancipatory research is to empower its participants and bring about a positive change for them, as opposed to for the benefit of researchers or institutions (as in traditional research) (Barnes & Sheldon 2007). Operating from an ‘emancipatory research’ standpoint, user involvement was central to the design of this project and its activities. The mentoring scheme and training was designed by people on the spectrum, including a member of the research team and members of an advisory panel.

The training was then evaluated through feedback forms given to mentors at the end of the mentoring session, which included both qualitative (open) and quantitative (closed) question formats. Further evidence on the relevance of the training was then reviewed following the delivery of the mentoring program through interviews conducted with the
participants. At the stage of the writing of this dissertation however, not all of this information had been collated, and so the data utilised for reflection in this dissertation have focused on the feedback gained from would-be mentors who undertook the training, whether they continued to be in the pool of mentors used for the full mentoring program or not.

3.2 Ethical considerations

The mentoring training and project as a whole was conducted according to the ethical guidelines of the British Education Research Association (BERA, 2011) for research with children and young people; and approved by a London South Bank University ethics panel.

Parameters for the role of the mentor were clearly defined in the training and guidance documents both for mentors and mentees for the full project. This included information on mentoring skills, risk assessment and safeguarding, and positive ways of working with autistic adults informed by an ‘insider perspective’. Following the training, all participants who wanted to become mentors needed to go through DBS checks and an interview process before being admitted to the mentor pool that was used for the full project.

All data from the project has been kept securely and anonymity of participants ensured. All participants will be debriefed about the findings of the project and a wider dissemination event has been planned for the full project.

Participants were invited to take part in the study through poster advertising displayed at a number of Universities, Further Education Colleges and public internet forums. The advert was also distributed by the National Association of Disability Practitioners (NADP). Respondents to the recruitment advert were then sent an introductory letter, participant
information sheet and expression of interest form (see Appendix 1). Consent was sought at
the beginning of this process and checked at intervals throughout the mentoring program.

During the process of recruitment it was decided to change the initial sampling of mentees
to be between the ages of 16-24 to be changed to 18+ in order to widen access to the
mentoring program.

3.3 Positionality statement

As a researcher who both has a diagnosis of Asperger’s and a father to an autistic son, I
bring with me an insider’s perspective to this study and to the mentoring project as a whole.
Whilst being a member of the autistic community may be said to bias me in favour of social
model approaches, this is reflected by many within the autistic community (Kenny et al.
2015; Milton, 2016a). The ethos of this project was mutually agreed upon by the other
researchers involved in the main mentoring project and the advisory group that was
established to advise the project. Indeed, having researchers from a multitude of
perspectives working on a project helped to maintain a participatory and emancipatory
ethos (Milton and Bracher, 2013).
Section 4: Research methods

In section 3 of this dissertation, the methodological rationale, ethical considerations and positionality of the researcher were explored. In section 4, the design of the mentor training is outlined, followed by a report on the piloting of the mentor training with the project advisory group. The design of the training feedback forms are then explained and a description of the sampling used on the project and proposed data collection for the mentoring project as a whole (this latter aspect feeding back into the development of the training materials).

4.1 The design of the mentor training

Following the remit of the mentoring project as a whole, the mentor training was constrained to one-day of training. Following the literature review outlined in section 2 of this dissertation and advice from the advisory group, the training day was broken up into thirty-minute time slots on a variety of topics as follows:

- An introduction to the project, aims of the training day, and introduction to mentoring.
- Autism in an historical and social context.
- A different way of thinking.
- Sensory perceptions and autism.
- Interaction and communication.
- Stress and anxiety.
- Autism and gender.
- The SPELL framework.
- Boundaries, recording risks and safeguarding.
- The Personal Well-being Index (PWI), goal-setting and Personal Construct Theory (PCT).
- Concluding guidance.

The introductory section of the training day sought to settle participants into the expectations of the day and the project, as well as introducing mentors to current theory and practice in the field of mentoring more generally. Within this section an exercise was designed for participants to reflect on the differences between mentoring and other support roles such as counselling, advocacy, befriending, and coaching.

A section was included on the historical and social context of autism, not only to set the scene as to where contemporary notions of autism originate, but to introduce models of disability and the framing of the project within a social model, emancipatory, and person-centred approach. This section is then followed by a number of core sections relating to theories of autism and aspects of the autistic experience. The section devoted to a ‘different way of thinking’ introduces participants to executive functioning theory, weak central coherence, and monotropism, as well as relevant criticisms of these theories and interactive activities and case study examples to help situate the theories in mentoring practice. A section was included regarding autism and gender following input from the advisory group and written and delivered by members of the advisory group.

The final aspects of the training day was to frame the ethos of the mentoring project, reviewing role boundaries, advising on safeguarding issues, and covering the data collection needed for the project. This included an introduction to the SPELL framework, as developed
by the National Autistic Society over many years (NAS, 2016b). This framework concentrates on the needs of autistic people in terms of: structure, positive approaches, (building) empathy, (employing) low-arousal (techniques), and (building) links (between the autistic person and significant others in their life).

4.2 Piloting the training with the advisory group

The training was initially piloted with the advisory group for the project. Feedback gained suggested that personal references in the training material needed to be taken out in order to standardise the information. It was also suggested that the training day needed less information and more time for discussion and activities related to the issues covered. The training was then adapted to take account of this feedback, personal references were removed, and information in many of the sections was made more succinct to add in more time for participant reflection, discussion, and example activities. The training was also edited following the experience of delivering subsequent training sessions. Four training days were conducted in the course of the project.

4.3 Design of training session feedback forms

Following the training sessions, participants were asked to rate the training session on Likert scales in the following categories, from strongly agreement to strong disagreement:

- A good balance of the information presented.
- Relevance and usefulness.
- Organisation and being easy to follow.
- The materials being useful.
- That the training would be helpful in their role as a mentor.
- That the trainers delivering the training were knowledgeable.

- That the training met expectations.

- The training being of the right length.

- The standard of the room and facilities within which the training took place.

Participants were also asked a number of qualitative questions regarding: what they liked most about the project, what could be improved in the training, the likely impact on their practice, if anything should be added or removed from the training, and whether they felt that they needed additional training in autism.

4.4 Design of the mentoring scheme

The full mentoring project looked to study the impact of access to mentoring on the wellbeing of twelve adults on the autism spectrum. Each participant received one hour of mentoring per week over a six month period. A period of six months was chosen because of the view expressed by adults on the autism spectrum in the Research Autism consultation held in 2007, that a short-term mentoring scheme would be most effective for them, and that they would not be looking for a long-term ‘befriending’ style of relationship, but short-term goal oriented support to help them move on with their lives.

A variety of mentoring arrangements were implemented in this project, including face-to-face interactions and email based interactions depending on the preferences of the participants. For some participants access to funded mentoring was possible. For example, some adults on the autism spectrum receive mentoring within the context of a package of support in College or University, or through an Access to Work grant. Where no such
arrangements were available the project recruited people who already had experience of supporting people on the autism spectrum. Mentors and mentees were matched as guided by the mentees’ goals for mentoring, which they identified on their expression of interest form/screening tool. All mentors and mentees were able to contact the research team during the programme in case of any issues that they feel unable to resolve within their mentoring sessions. Additionally, mentors were invited to attend a peer support session approximately three months into the mentoring programme.

4.5 Sampling

Opportunity sampling was utilised to recruit both mentors to the training program and for mentees. Mentors were recruited through advertising (see section 3.2) at a number of organisations and internet forums. Those who received the training then were invited to apply to the mentor pool for the main project.

4.6 Data collection and analysis for main mentoring program

The research team employed the standardised tool, the PWI-A (2006) developed by the International Wellbeing Group at Deakin University in Melbourne, to measure changes in participants’ perception of their own wellbeing prior to, and after completion, on the mentoring programme. This tool asks participants to rate their standard of living, personal health, achievement in life, personal relationships, personal safety, community connectedness, future security, and spirituality and religion.

This data was supported by qualitative indicators from a number of data collection methods. Semi-structured interviews were conducted with both mentors and mentees after completion of the mentoring programme. The mentors were also given materials
based on the Salmon Line technique (as used in PCT, Salmon 2003) to utilise with participants through the mentoring programme that were also used to analyse progression from the subjective viewpoint of the participants (see Appendix 2). Additionally, mentors and mentees were asked to complete a mentoring record sheet and reflective journal after each mentoring session. Qualitative data from this is now being analysed using thematic analysis (Braun & Clarke 2006). Questions were included in the interviews with mentors following the program that were specifically focused to feedback into the development of the training materials used in the project.
Section 5: Results and analysis

The training feedback forms used in this project used both closed and open-ended questions to produce quantitative and qualitative data to help answer the following research questions:

What information and activities are most effective for meeting the training needs of would-be mentors of autistic adults?

What aspects of a training programme for would-be mentors of autistic adults are most appreciated by those undertaking the training?

In this section results from the four training sessions conducted are reported on. Across these four sessions, 45 participants agreed to fill out these feedback forms.

5.1 Analysis of training feedback – quantitative data

Responses to the Likert-scale questions contained in the training feedback forms are shown below for the four training sessions conducted (SA = Strongly Agree, A = Agree, N = Neither agree nor disagree, D = Disagree, SD = Strongly Disagree; average score was found by scoring each ranking 5, 4, 3, 2, and 1 respectively):
### Table 5.1.1 Feedback from training day one:

<table>
<thead>
<tr>
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<th>SA</th>
<th>A</th>
<th>N</th>
<th>D</th>
<th>SD</th>
<th>No Answer</th>
<th>Avg. score</th>
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<td>-</td>
<td>-</td>
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<td>-</td>
<td>-</td>
<td>-</td>
<td>4.88</td>
</tr>
<tr>
<td>Met expectations</td>
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<td>7</td>
<td>-</td>
<td>2</td>
<td>-</td>
<td>-</td>
<td>4.24</td>
</tr>
<tr>
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<td>-</td>
<td>-</td>
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<td>8.5</td>
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<td>1</td>
<td>-</td>
<td>3.38</td>
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</table>

### Table 5.1.2 Feedback from training day two:

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<th>N</th>
<th>D</th>
<th>SD</th>
<th>No Answer</th>
<th>Avg. score</th>
</tr>
</thead>
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<td>-</td>
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<td>-</td>
<td>1</td>
<td>-</td>
<td>4.23</td>
</tr>
<tr>
<td>Organised/easy to follow</td>
<td>5</td>
<td>4</td>
<td>3</td>
<td>1</td>
<td>-</td>
<td>-</td>
<td>4</td>
</tr>
<tr>
<td>Materials useful</td>
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<td>6</td>
<td>1</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>4.38</td>
</tr>
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<td>Help in role as mentor</td>
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<td>1</td>
<td>-</td>
<td>1</td>
<td>4</td>
</tr>
<tr>
<td>Trainers knowledgeable</td>
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<td>1</td>
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<td>-</td>
<td>1</td>
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<td>4</td>
</tr>
<tr>
<td>Right length</td>
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<td>2</td>
<td>2</td>
<td>1</td>
<td>-</td>
<td>3.62</td>
</tr>
<tr>
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<td>6</td>
<td>1</td>
<td>-</td>
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Table 5.1.3 Feedback from training day three:

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<th>N</th>
<th>D</th>
<th>SD</th>
<th>No Answer</th>
<th>Avg. score</th>
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<td>-</td>
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<td>-</td>
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<td>3</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>4.5</td>
</tr>
<tr>
<td>Organised/easy to follow</td>
<td>3</td>
<td>3</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>4.5</td>
</tr>
<tr>
<td>Materials useful</td>
<td>2</td>
<td>4</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>4.33</td>
</tr>
<tr>
<td>Help in role as mentor</td>
<td>2</td>
<td>3.5</td>
<td>0.5</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>4.25</td>
</tr>
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<td>Trainers knowledgeable</td>
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<td>1</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>4.83</td>
</tr>
<tr>
<td>Met expectations</td>
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<td>4</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>4.33</td>
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<td>-</td>
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<td>-</td>
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</tr>
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Table 5.1.4 Feedback from training day four:

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<tr>
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<th>N</th>
<th>D</th>
<th>SD</th>
<th>No Answer</th>
<th>Avg. score</th>
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<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>4.63</td>
</tr>
<tr>
<td>Relevance/useful</td>
<td>7</td>
<td>1</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>4.88</td>
</tr>
<tr>
<td>Organised/easy to follow</td>
<td>5</td>
<td>3</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>4.63</td>
</tr>
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<td>4.75</td>
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<td>-</td>
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<td>-</td>
<td>4.88</td>
</tr>
<tr>
<td>Trainers knowledgeable</td>
<td>8</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>5</td>
</tr>
<tr>
<td>Met expectations</td>
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</tr>
<tr>
<td>Room/facilities</td>
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<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>4.88</td>
</tr>
</tbody>
</table>
Feedback from the initial training days was generally very positive, particularly in regard to the knowledge of the trainers delivering the programme, but also in terms of the training being easy to follow, well organised, and with useful materials. A similar pattern was found with the last two training days, with the knowledge of the trainers being the most highly scored. The final training day was scored the highest of the four, indicating that the small changes made during the process and the trainers becoming more acquainted with the materials and timings, were beneficial to participant perceptions of the training program. A comparison of the training day feedback can be seen in the table below:

Table 5.1.5 Collated feedback from all four training days:

<table>
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<th>T2</th>
<th>T3</th>
<th>T4</th>
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<td>Relevance/useful</td>
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<td>4.5</td>
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<td>4</td>
<td>4.33</td>
<td>4.5</td>
</tr>
<tr>
<td>Right length</td>
<td>4.41</td>
<td>3.62</td>
<td>4.33</td>
<td>4.5</td>
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<tr>
<td>Room/facilities</td>
<td>3.38</td>
<td>4.38</td>
<td>4</td>
<td>4.88</td>
</tr>
</tbody>
</table>

The lowest marks were given during the second and third training days in regard to the training day being of the right length and balance. This could indicate that a longer period of training would be ideal. It should be noted that the only negative comments about the training came from a couple of participants who attended the second and third training
days. Of the others who attended and offered feedback, the vast number of responses were very positive and encouraging.

5.2 Analysis of training feedback – qualitative data

Qualitative data was collated in regard to a number of questions (as outlined in section 4) and then thematically categorised. In terms of what participants liked most about the training the main themes were as follows (in order of the occasions mentioned by participants throughout the four training sessions):

- Speakers being on the autism spectrum

“Valuable to have personal experiences and interpretations of the autism spectrum.” (Participant from training day 2).

“The trainers talking about their experiences.” (Participant from training day 2).

“Input from trainers with personal experience of autism.” (Participant from training day 3).

The most often aspect of the training that attendees most appreciated was having the training delivered with significant input from more than one autistic trainer.

- The balance, variety and structure of the content

“The structure meant that we could properly engage with the content – clear sections split with tasks, lectures and breaks accordingly.” (Participant from training day 1).

“There was a good balance between the lectures and interactive exercises, which I enjoyed.” (Participant from training day 4).

The breaking down of the training into sections that allowed participants to apply their learning to case examples was also highly appreciated by participants.
- **Activities and discussions regarding strategies**

“Being in a group and question / sharing with peers and trainers.” (Participant from training day 3).

A number of participants commented on the ability to discuss and question issues in-depth with the other participants and trainers.

- **Straightforward and informative materials**

“Straightforward and very informative.” (Participant from training day 2).

“Having a specific focus for everything we have learned.” (Participant from training day 3).

- **The talk on autism and gender being included**

“The section on...girls with autism (very rare for such things to be mentioned.” (Participant from training day 4).

“Also, the attention to how female individuals and gender differences may manifest in autism.” (Participant from training day 3).

The section of autism and gender, delivered by a member of the advisory group for the mentoring project was often highlighted as a particularly strong aspect of the training.

- **Networking and learning from others**

“Meeting others and listening to the presentations.” (Participant from training day 4).

- **Focus on more effective relationships**

“Focus on how to create more effective relationships and communication with autistic learners.” (Participant from training day 2).

- **Less deficit-driven and mention of rarely covered issues (such as ‘shutdowns’)**
“Less deficit-driven.” (Participant from training day 4).

The social model and person-centred account given to frame the project was also well received by participants.

The best aspect of the training in the perception of participants was that the training was delivered by people with personal experience of autism. This was closely followed by the balance, variety and structure of the content, and the interactive activities of the training program. The aspect of the training program designed and delivered by members of the advisory group regarding autism and gender was also very well received. Only one negative or critical response was found in regard to the initial question of what participants liked most about the training. This comment suggested that the training was ‘not fit for purpose’, as it did not cover in their view the differences between mentoring autistic and ‘NT’ people.

The second open question was in regard to what could be improved about the training. The main themes found in response to this question were:

- More interactive activities or videos of mentoring in practice

“More specific and 1-2-1 exercises.” (Participant from training day 2).

“Perhaps role-playing / videos of mentoring in action.” (Participant from training day 3).

“A common response to how the training could have been improved was to have included yet more interactive exercises and also video-clips or role-play activities.

- Perhaps delivered over two days / too much information in time available
“Possibly tried to cover too much in the time available (but all content was relevant).”  
(Participant from training day 1).

“Perhaps the training could be delivered over two days.”  
(Participant from training day 1).

Another very common response was to expand the training over a two-day period in order to fit in more discussion and activities.

- More information and advice on mentoring in practice and practical strategies

“Focus more on strategies as a mentor, rather than ‘what is autism?’”  
(Participant from training day 1).

“More on mentoring in practice.”  
(Participant from training day 1).

- Environmental factors about the venue

“Chairs facing the front.”  
(Participant from training day 2).

- Participants not wanting to change anything about the training

“I found it interesting and at the right level.”  
(Participant from training day 4).

“I don’t think it could have been improved.”  
(Participant from training day 4).

- The pace of the training – for some too slow, others too fast

These responses indicate that the training was generally well received, yet would have benefitted with yet more time for interactive activities, discussion of practical strategies, and delivered over a longer period of time.
The third open-ended question regarded what impact the training would have on the practice of the participants. The following main themes were found:

- Increasing understanding in person-centred way

“I feel I have a further understanding of autism.” (Participant from training day 1).

“More tools for the toolbox and increased understanding.” (Participant from training day 2).

“Beware of assumptions.” (Participant from training day 3).

“Help with understanding autism, and the promotion of a person-centred way of working.” (Participant from training day 4).

- The use of PCT and the Salmon line

“Shall incorporate the Salmon line – thank you.” (Participant from training day 2).

“Liked the Salmon line.” (Participant from training day 3).

“PCT and the Salmon line – a useful visual support.” (Participant from training day 4).

Following general comments regarding an increased understanding, the Salmon line tool was highlighted by a high number of participants as having influenced them regarding their future practice.

- Providing structure and practical strategies

“The SPELL framework was new to me and may influence my practice as a mentor.” (Participant from training day 2).

“It has clarified where the boundaries lie and it is good to have the structure to the sessions.” (Participant from training day 3).
- Feeling empowered as practitioners

“Feel more empowered.” (Participant from training day 4).

- Would have little impact on practice

“Remember to check out the mentee’s reasons for stuff.” (Participant from training day 2).

It is clear from these responses that the person-centred ethos coupled with the practical strategies and materials offered by the project were seen as useful by the participants, particularly the use of the Salmon line. Only two participants commented that they thought that the training would have little impact on their practice (from 45 respondents).

When asked if participants would change, add or remove anything from the training program, the following themes were identified:

- Not changing anything

“N/A.” or “No.” (Participant(s) from all training days).

- More on mentoring case studies and examples

“More on the mentor role.” (Participant from training day 1).

“More case studies.” (Participant from training day 1).

- More exercises and activities

“More in the way of practical strategies.” (Participant from training day 4).

- Some slides could be delivered as a handout

“The last five slides could have been given as a handout.” (Participant from training day 3).

- Less on autism history

- Creating videos of those who have experienced mentoring
By far the most common response to this question was that the training should not be changed at all. However, some participants wanted to have more time with interactive activities and be given more practical examples. These issues could be addressed if the training were to be expanded into a two-day model.

Lastly, when asked what additional training the participants felt they needed, the most common themes were:

- A simple response suggesting additional training would be always helpful
  “May do further on the journey of mentoring. Always learning.” (Participant from training day 4).

- A meeting once/twice a year to refresh knowledge
  “A training / meeting once or twice a year at least – to refresh and facilitate good practice.” (Participant from training day 1).

- Videos of good / bad mentoring sessions
  “A video showing a good and a bad mentoring session.” (Participant from training day 3).

- Looking into overlaps with mental health
  “Overlaps when working with mental health and an autism diagnosis.” (Participant from training day 1).

- Navigating forms and bureaucracy
  “How to navigate bureaucracy.” (Participant from training day 4).

- Not needed
Only one participant stated that they did not need additional training. Some wanted more information on the overlap with the experience of mental health difficulties. The suggestion of meeting again to refresh knowledge however directly influenced the structure of the main mentoring program.

5.3 Initial findings from mentoring program

The initial findings that are currently being collated for the mentoring project have been very encouraging, with every mentee that has finished the full six-month program showing increases in well-being and substantial progress toward their self-selected goals. When asking mentors to look back on hindsight on the training, responses again have been very positive about the usefulness of it, and the benefit of having training designed and delivered with significant input from autistic people. The use of the Salmon Line was also referenced as being of particular practical usefulness. Again though, mentors described how the training could have been perhaps better delivered over two days instead of one. On mentor suggested that they had not gained a great deal from the section of the training regarding the SPELL framework, however this comment came from a participant who already had a great deal of experience in the field and with mentoring.

During the mentoring program it was found that the reliability of mentors was variable, and seemed to have an impact on the quality of the mentoring relationship. Although consistency and reliability was emphasised in the mentoring training, this may show that this aspect needed added emphasis. Similarly, in regard to advice for mentors to facilitate the initial setting of goals, mentees sometimes struggled with this aspect of the program at first, and some mentors struggled to facilitate this process.
Section 6: Conclusions

This study looked to answer the following research questions:

What information and activities are most effective for meeting the training needs of would-be mentors of autistic adults?

What aspects of a training programme for would-be mentors of autistic adults are most appreciated by those undertaking the training?

In section 5: results and analysis, it was shown that the training day was well received by the vast majority of participants. Those who also went on to become mentors on the project, found the advice and materials offered of practical use, particularly in regard to the Salmon line tool. The strongest aspect of the training was that it was designed and delivered with significant autistic input. Tackling issues from a social model and ‘insider’ perspective was greatly appreciated by participants. The main criticisms of the training regarding its length and having more time to explore the issues at hand, with more practical examples and strategies. Some participants reflected upon how the use of video clips of mentoring in progress would have enhanced the training. The most common suggestion from participants to resolve these issues was to extend the training over two days.

In this section, these findings will be discussed in more depth, and limitations of the study investigated. This section will conclude by exploring possible next steps for the mentoring project as a whole, and reflect upon how the training aspect of the project can potentially be improved.
6.1 Training being informed by the authentic voice of autistic people

Throughout the four training sessions that were completed, participants most strongly referred to having the authentic voice of autistic people delivering the content. Autistic input in the design and delivery of the training significantly affected the content of the materials as well as allowing for nuanced explanations and examples presented and answers given to participant questions. A number of the participants also commented favourably on the inclusion of autism and gender as a topic and more than one autistic trainer with differing dispositions and experiences.

This finding was further highlighted by the quantitative data gathered from the participant feedback forms following the training. The trainers being knowledgeable about the information being presented was consistently scored highest in all of the four training sessions. It may be the case that this was due in part to the specific personal qualities of the people (both autistic and non-autistic) who delivered the training. If this training model were to be expanded and evaluated on a wider scale however, one would be able to look into more depth at these issues. It is a certainly a recommendation of this project however, that due consideration is given to the participation of autistic voices in the development of training materials.

The majority of work that has previously been undertaken in the area of mentoring autistic people has been situated within a medical / behavioural model and also designed without significant autistic input (Griffith et al. 2012; Curtin et al. 2015; Arnes et al. 2015), and few have been subject to rigorous evaluation (Gelbar et al. 2013). This has led to the focus of such programs being primarily aimed at concerns driven by the non-autistic population, such as the learning of ‘social skills’ (Milton, 2016a) and often normative, remedial and
normalising in outlook. Current mentoring practice for autistic people in the UK often is not supported by adequate training, or training at all, yet recent work by Hamilton et al. (2016) suggest such training is essential for successful mentoring practice. This project has given further evidence that such training can have added impact by being led by the concerns of autistic people, and following a person-centred and participant-led model.

6.2 Developing the training program into a two-day model

The feedback from participants to the mentoring training, alongside the initial findings from the full mentoring project, have both been very encouraging. Whilst aspects of the training have been highlighted that could be improved, for example: more emphasis on reliability and consistency, and the setting of goals, by far and away the most significant aspect of the training that could be improved upon was that of exploring how the training could be expanded into a two-day model. From the outset of the project however, this was recognised as a potential issue, as it was not an easy process to cover so many significant issues within one day of training. For this reason, the training was developed to be deliverable in 30 minute sections. Given that there had been a comment regarding the need for the SPELL framework within the training, as well as positive comments on this aspect, it could be the case that this could be repositioned within the training schedule, or more time given to giving practical examples and discussion of the framework. If the training was to be delivered over a two-day period, the first day could concentrate on familiarising participants with the project and core aspects of autism, whilst the second day could concentrate on mentoring practice utilising what had been learnt on the first.
Day one:

- An introduction to the project, aims of the training day, and introduction to mentoring.
- Autism in an historical and social context.
- A different way of thinking.
- Sensory perceptions and autism.
- Interaction and communication.
- Stress and anxiety.
- Autism and gender.

Day two:

- The SPELL framework.
- *Applying theory in context – autistic ways of perceiving and thinking.*
- *Applying theory in context – interaction and communication.*
- *Applying theory in context – stress and anxiety.*
- Boundaries, recording risks and safeguarding.
- The Personal Well-being Index (PWI), goal-setting and Personal Construct Theory (PCT).
- Concluding guidance.
The above list includes in italics possible sections that could be added if the training was to be delivered over two days, allowing for greater time for participants to reflect, discuss and apply theory within the mentoring context.

Whilst it would have been of great benefit to have produced resources such as video footage of mentoring and case studies, this was beyond the remit and budget of the project. It is certainly the case however, that such activities in future could enhance the mentoring training.

6.3 Limitations of the study

Funding was fairly small for the project, and thus additional learning materials such as video clips could not be commissioned. This was highlighted by participants who undertook the training, but could be seen as a future project in and of itself.

The training events were perhaps attractive to participants as they were offered for free, with many not applying to become part of the wider mentoring project. However, there participation in the evaluation of the training was helpful and the training may have helped them in whatever settings they work in and when they had future interactions with autistic people.

The success of the training could have been due to the personal qualities of the trainers, yet an expanded research program could explore this issue in more depth.

6.4 Next steps for the development of a mentoring training program

In order for the project to realise its emancipatory aim of ultimately producing an intervention informed by, and practically useful, to autistic people the project could be
expanded by looking to accredit the training program. If more funding can be secured to extend the project, the training could be developed into a two-day model. Trainers could be identified and trained, and then evaluation of the model undertaken on a wider scale.

6.5 Key findings and recommendations

The findings from evaluating the feedback forms following the training sessions, as well as initial findings and feedback from mentors who continued on to the mentoring program, show clearly the value of training that has had significant input from autistic people in its design and delivery. The framing of the training within a social model, participatory and person-centred framework was also well received by participants and practical tools such as the Salmon line found to be of particular use. Participants felt that they had benefitted from the face-to-face and interactive nature of the training, and the quality of materials they received. Ideally however, the training would be extended over a two-day period. The initial findings from the full mentoring program suggest that extra emphasis may be needed in the training also on issues such as the reliability and consistency of mentors and mentoring arrangements, and the facilitating of setting goals and role boundaries. These issues could be addressed in more depth, as well as more interactive activities added, if the training was to be delivered over a longer period of time. This would also answer the main criticisms received from participants that the training may not have been of the right length or balance (for some participants).

Through the literature review, it was found that current practices are not supported by a strong evidence base, and that improvements to training, supervision and CPD of mentors working with autistic people is needed (Hamilton et al. 2016). Given the positive findings from this project and the training feedback received, it is proposed that a larger scale
project is required to build upon the model developed in this project, with the view of operationalising a mentor scheme that includes the significant components so far identified.

6.6 Concluding remarks

As the full mentoring project progresses, it is becoming clear that there are many potential benefits to a time limited and goal oriented mentoring model, when founded upon an ethos, training, and supervision model that has had significant input from autistic people. There has been a lack of rigorous evidence to base current mentoring practice with autistic adults upon, and much of the current evidence has been based on normative / deficit models with little autistic participation into the design and delivery of training programs. To offer mentoring to autistic people without a formalised and accredited training process and supervision structure is potentially damaging. Specific training in autism, let alone incorporating a social model perspective, person-centred approaches and insider viewpoints, does not seem to be currently required for practitioners to work as mentors with autistic adults, nor are there essential safeguards for both mentees and mentors.

The support needs of autistic adults can be very varied and idiosyncratic, but could include anything from managing finances and practical everyday living, to accessing education, training and employment. The needs of autistic adults are often unrecognised however, particularly when one takes into account the mutual incomprehension that ensues from the double empathy problem (Milton, 2012a; 2014a; Chown, 2014). Autistic adults have reported how current models of support have not met their needs, whilst mentoring can help with their progression and meeting goals. This project has shown that the input of well-qualified autistic people in the design and delivery of training can significantly enhance the
experience and understanding gained by participants, a finding which has ramifications beyond that of mentoring practice.

(16,381 words)
Bibliography


National Autistic Society (2016b) SPELL [online]:


Appendices

Appendix 1: Introductory letter

Participant Information Sheet: Mentors

Research Autism Cygnet Mentoring Project

You are being invited to take part in a research study. Before you decide it is important for you to understand why the research is being done and what it will involve. Please take time to read the following information carefully. Talk to others about the study if you wish.

Ask us if there is anything that is not clear or if you would like more information. Take time to decide whether or not you wish to take part.

During the Collaborative Autism Research Forum (CARF) in 2009, adults with autism recommended that Research Autism should develop a specialised mentoring or coaching programme for young adults on the autism spectrum. Much mentoring that already exists is not “autism-friendly” or equates more closely with befriending, which is something adults with autism told CARF they did not find helpful. There is also a lack of evaluation of existing mentoring schemes.

The aim of this study is to develop a specially designed mentoring programme for adults on the autism spectrum and to assess its effectiveness.

You have been invited to participate in this study as you have expressed an interest in mentoring a young person on the autism spectrum.

In total, 24 participants will be included in the study. 12 of these will be mentors and 12 will be people who will receive mentoring.

It is up to you to decide whether or not to take part. If you do, you will be given this information sheet to keep and be asked to sign a consent form. You are still free to withdraw anytime during the project without giving a reason. A decision to withdraw, or a decision not to take part, will not affect any mentoring or support you are already in receipt of.

If you are willing to participate, you will be matched to a suitable mentee who will be identified based on matching their goals for mentoring to your skills and experience. Prior to starting the mentoring, you will be asked to complete a standardised questionnaire which assesses Quality of Life (QoL). QoL can refer to aspects of a person’s well-being (physical, psychological, social), as well as aspects of the environment.

You will then be invited to come to London Southbank University (or an alternative place if more convenient) to meet with your mentee and a research assistant. The research assistant will introduce you to your mentee, briefly explain to both of you the ground rules for the mentoring relationship. You will then assist your mentee to devise (up to three) goals for mentoring and decide how (face-face, telephone, on-line), when and where mentoring will occur. Mentoring will last for 6 months and occur for an hour each time on a weekly basis (maximum). Throughout the mentoring
you will both be asked to keep brief recordings of your meetings. In the last mentoring session, you will be asked to facilitate the mentee in reviewing the goals set during the first session to see whether things have changed. You will also be asked to complete a brief form to assess whether your own goals for participating in the mentoring programme were addressed. At the end of the mentoring (6 months), you will be asked to complete the same standardised QoL questionnaire which you completed at the start of the study. You will also be asked to participate in an interview to explore your experiences of participating in the programme. The interview will last for approximately one hour and will be audio recorded. If you prefer, the interview can be carried out via telephone or online. 6 months after the mentoring has ended you will be asked to complete the standardised QoL questionnaire one final time.

It is not anticipated that you will be disadvantaged or suffer any risk form this study. However, some issues that your mentee chooses to discuss with you may be difficult or upsetting to talk about. You will be offered regular supervision to help you manage any distressing experiences. If your mentee discloses anything which suggests your mentee or someone else could be at risk of harm you will be required to report it to a research assistant on the project.

It is not known whether you will benefit directly from being a mentor, however the following have been suggested as potential benefits (for mentors) of mentoring:

- improved emotional intelligence skills and capabilities;
- Knowledge and skill development:

Your participation will be invaluable in helping us to develop an appropriate mentoring approach for people on the autism spectrum.

You are free to withdraw from the study and not have your information included, at any time up to the completion of the project report or publications. After that time, it would be impossible for the researcher to remove the information you provided.

All information received from you will be handled in a confidential manner and stored in a locked filing cabinet and on a password protected computer in an environment locked when not occupied. Only the researchers working on the study will have direct access to the information. Any reference to you will be coded. This information will be held until September 2016.

This study is being completed at London South Bank University and funded by Research Autism. It has been reviewed and ethically approved by the London Southbank University Research Ethics Committee.

If you have a concern about any aspect of this study, you should ask to speak with the research assistants who will do their best to answer your questions (Damian Milton [miltond@lsbu.ac.uk] and Tara Sims [simst@lsbu.ac.uk]). If you wish any further information regarding this study or have any complaints about the way you have been dealt with during the study or other concerns you can contact Dr Nicola Martin (martinn4@lsbu.ac.uk), who is the Principal Investigator for this study. Finally, if you remain unhappy and wish to complain formally, you can contact the Chair of the University Research Ethics Committee. Details can be obtained from the university website:

https://my.lsbu.ac.uk/page/research-degrees-ethics

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Participant Information Sheet: Mentees

Research Autism Cygnet Mentoring Project

You are being invited to take part in a research study. Before you decide it is important for you to understand why the research is being done and what it will involve. Please take time to read the following information carefully. Talk to others about the study if you wish.

Ask us if there is anything that is not clear or if you would like more information. Take time to decide whether or not you wish to take part.

During the Collaborative Autism Research Forum in 2009, adults with autism recommended that Research Autism should develop a specialised mentoring or coaching programme for young adults on the autism spectrum. Much mentoring that already exists is not “autism-friendly” or equates more closely with befriending, which is something adults with autism told CARF they did not find helpful. There is also a lack of evaluation of existing mentoring schemes.

The aim of this study is to develop a specially designed mentoring programme for adults on the autism spectrum and to assess its effectiveness.

You have been invited to participate in this study as you are a young adult who identifies with autism and you have identified that mentoring may be helpful to you.

In total, 24 participants will be included in the study. 12 of these will be mentors and 12 will be people who will receive mentoring.

It is up to you to decide whether or not to take part. If you do, you will be given this information sheet to keep and be asked to sign a consent form. You are still free to withdraw anytime during the project without giving a reason. A decision to withdraw, or a decision not to take part, will not affect any mentoring or support you are already in receipt of.

If you are willing to participate, you will be matched to a suitable mentor who will be identified based on your goals for mentoring. Prior to starting the mentoring, you will be asked to complete a standardised questionnaire which assesses Quality of Life (QoL). QoL can refer to aspects of a person’s well-being (physical, psychological, social), as well as aspects of the environment (Michielson et al 2011)

You will then be invited to come to London Southbank University (or an alternative place if more convenient) to meet with your mentor and a research assistant. The research assistant will introduce you to your mentor, briefly explain to both you and your mentor the ground rules for your mentoring relationship. You and your mentor will then collaboratively devise goals for your mentoring and decide how (face-face, telephone, on-line), when and where your mentoring will occur. Mentoring will last for 6 months and occur for an hour each time on a weekly basis (maximum). Throughout the mentoring you and your mentor will be asked to keep brief recordings of your meetings. In the last mentoring session, you will be asked to review the goals you set during the first session to see whether things have changed for you. At the end of the mentoring (6 months), you will be asked to complete the same standardised QoL questionnaire which you completed at the start of the study. You will also be asked to participate in an interview to explore your experiences of receiving
mentoring. The interview will last for approximately one hour and will be audio recorded. If you prefer, the interview can be carried out via telephone or online. 6 months after the mentoring has ended you will be asked to complete the standardised QoL questionnaire one final time.

It is not anticipated that you will be disadvantaged or suffer any risk from this study. However, some issues you discuss with your mentor may be difficult or upsetting to talk about. It is up to you what you wish to share and discuss with your mentor. If you discuss anything which your mentor feels suggests you or someone else could be at risk of harm they will be required to report it to the research assistants and you may be referred to other services or safeguarding procedures may need to be followed.

It is hoped that you will benefit from receiving mentoring, however we cannot confirm this as this is part of what the project will be evaluating. Your participation will be invaluable in helping us to develop an appropriate mentoring approach for people on the autism spectrum.

You are free to withdraw from the study and not have your information included, at any time up to the completion of the project report or publications. After that time, it would be impossible for the researcher to remove the information you provided.

All information received from you will be handled in a confidential manner and stored in a locked filing cabinet and on a password protected computer in an environment locked when not occupied. Only the researchers working on the study will have direct access to the information. Any reference to you will be coded. This information will be held until September 2016.

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https://my.lsbu.ac.uk/page/research-degrees-ethics
Appendix 2: The Salmon line tool

On the lines below, please mark how satisfied you are with your current level of attainment of each of your goals for mentoring

Goal 1: ________________________________________________________________

____________________________________________________________

1  2  3  4  5  6  7  8  9  10

(Adapted from Salmon 1988*)

The person is asked to place him/herself on the line in relation to their perceptions of how they are performing in this area currently. Ideas for progressing towards achievement can be generated in small, achievable steps.

For each aspect of life that is chosen, it is represented on a continuum with one end representing satisfaction in that area and the other representing dissatisfaction.