Scottish Borders
Autism Strategy

Autism and me
CONTENTS

1. Foreword P.3
2. Our Vision P.4
3. What is Autism? P.6
5. The National Picture of Autism P.9
6. The Local Picture of Autism P.16
7. Autism, in Numbers P.20
8. Towards an Autism Strategy for the Borders P.22
9. What is Working Well? P.25
10. Priority Areas Identified for the Borders P.26
11. Next Steps P.38

Appendices

Appendix A: List of previous autism initiatives, cited in the Scottish Strategy for Autism. P.39
Appendix B: Breakdown of current estimated spend on people with autism. P.40
Appendix C: Full list of services specifically named by questionnaire respondents as in need of further autism training. P.41

References P.42
Glossary P.44
The Scottish Strategy for Autism was published in 2011; marking growing recognition that autism deserves an agenda in its own right. The development of the national strategy ensures progress will continue to be made in delivering quality services for individuals on the spectrum.

In response to the national agenda for autism, the Scottish Borders has developed a local strategy, to ensure the needs of individuals on the spectrum of all ages living in the Borders will be acknowledged, and addressed.

A group of professionals from the National Health Service (NHS), education, social work and the third sector, including parents and carers, have been meeting to develop this strategy. The group has discussed the priority areas of need, identified through the Autism Mapping Project, conducted in 2013, and through professional and personal contributions.

The core part of the strategy is split into seven main sections addressing each of the priority areas, each linked to a section in the delivery plan. The thematic sections detail local context, a strategic overview and findings from our engagement process (including the mapping project).

Seven priority areas have been identified for action, locally:

1. **Autism awareness and training** – Improving public and professional awareness and understanding of autism;

2. **Diagnosis** – Good-quality early diagnosis and intervention for both children and adults;

3. **Getting the right services at the right time, for adults with autism and no learning disability** – Addressing the eligibility criteria and improving access to appropriate support;

4. **Purposeful occupational activities** – Finding the right opportunities to pursue individual interests and employment;

5. **Social support and opportunities** – Opportunities to develop social skills and access a range of social opportunities;

6. **Improving access and provision of housing** – Addressing the barriers to finding, securing and maintaining individual housing needs;

7. **Ensuring inclusion for people with autism and their families** – People will have their voices heard and acknowledged by professionals.
OUR VISION

It is our vision that people with autism and their families, living in the Scottish Borders, feel accepted and valued by their community, and have equal access to knowledgeable services, when they need them, so that they are able to live the lives they choose.

Our vision symbolises the fundamental principles we will aspire to, over the next ten years and beyond and has been shaped entirely by people with autism, families and professionals from a range of backgrounds.

How will we achieve our vision?

The purpose of this strategy is to help guide us towards realising this vision, by setting the tone and direction for improving outcomes for people with autism and their families, across the lifespan. Through our engagement, we have learned what is important to people living in the Borders and what they think needs to change. Our vision exists to represent these changes and outcomes.

The delivery plan sets out some of the actions under the priority areas in need of change, to help us achieve our vision. The delivery plan sets out these changes over the next 2, 5, and 10 years and incorporates the feedback we have received through our engagement.

The services and support referenced throughout this strategy and plan will be provided by the Scottish Borders Council, NHS Borders and our partners (*).
WHAT IS AUTISM?

Autism is a complex, lifelong developmental disorder and is sometimes referred to as autism spectrum disorder (ASD) or autism spectrum condition (ASC).

Autism is known as a spectrum condition, incorporating a wide range of characteristics that some people may or may not share, which can make some aspects of life difficult. Just like the non-autistic population, people with autism can have any measure of intelligence (IQ); however, people diagnosed with Asperger Syndrome or High Functioning Autism will typically have an average or greater IQ.

Therefore, each individual with autism is unique and may or may not require support, in various forms, to live an independent life.

There are broad characteristics that are common to all individuals on the autism spectrum, which can be present in varying degrees of severity. These are sometimes referred to as the ‘triad of impairments’ and currently form the basis of the diagnostic assessment for autism.

These common characteristics are detailed below, along with more specific details of difficulties people with autism may experience:

**Social communication (i.e. Use of verbal and non-verbal language)**

- Language processing
- Varying speech abilities
- Understanding language
- Using and understanding appropriate facial expressions, eye contact and body language

**Social interaction**

- Working out other people’s thoughts, emotions and expectations
- Picking up on unwritten social rules
- Social skills
- Building and maintaining relationships

**Social Imagination/Flexibility of Thought**

- Need for structure and routine
- Self-organisation - the concept of time and planning ahead
- Coping with change
- Transferring skills and knowledge from one situation or task to another

(Note, the American Psychiatric Association classification of autism has recently changed)
Perceiving the world and other people differently

Autism can be defined as a distinct cognitive style, or way of thinking, according to the ‘social model’ of autism. Therefore, people with autism interpret and process information about the world and other people differently. Difficulties arise due to attitudes and structure in society, leading to a misunderstanding of this thinking style. This is in contrast to the ‘medical model’ of autism, which classifies autism as a disorder; limiting one’s ability to function.

It is important to note that a range of additional characteristics have become associated with autism; some people with autism, but not all, may experience them. Some of these characteristics are listed below, with further detail provided for your information.

Stress and anxiety

It is generally accepted that people with autism often have associated anxiety and stress issues. Anxiety and stress can happen for a number of reasons, and people with autism can vary in their ability to manage these emotions. This may be because of difficulties with a lack of predictability and control, relating to others, and other factors which affect the individual’s sense of stability.

Over- or under-sensitivity to noise, lights, textures, balance, taste and/or proprioception (sensing your own body in relation to the physical space)

Some people with autism experience an over- or under-sensitivity to sensory stimuli, present in the environment. Over-sensitivity to stimuli can be overwhelming and potentially painful for people with autism. Loud, or unexpected noises can cause significant stress for the person. Under-sensitivity to stimuli can be potentially harmful too, with people unaware of injuries they have sustained, or not picking up on incoming information or messages being sent through the senses.

Co-existing conditions

People with autism may also present with characteristics of other difficulties/diagnoses, such as: ADHD, dyslexia, dyspraxia, dyscalculia, depression, anxiety, eating conditions and sleep disorders.

Use of language in our strategy

The complex nature of autism gives rise to a number of personal and professional perspectives, therefore it can be difficult to find a common language to use. However, it is important to note that we will adopt the language and terminology used within the National Autism Strategy; as such, we have tried to reflect the diversity of this community in a positive way.

Furthermore, we understand there is a need to be sensitive when using words like ‘impairment’ or ‘disorder’. These words are recognised clinically and in a professional realm, however many people with autism do not accept these terms, instead preferring to stress that they have a different way of perceiving and engaging with the world.

We have used the term ‘autism’ throughout our strategy to represent the whole spectrum; this includes Asperger Syndrome, High Functioning Autism, Atypical autism and Pervasive Developmental Disorder Not Otherwise Specified.

For a full list terms and descriptions, please see the Glossary of terms.
It is also important to understand that no two people with autism are alike, therefore each individual’s needs are unique and personal. Some people with autism may require a high level of support, on a daily basis, to help them with daily tasks, while others may only need a little support now and again to help them when they need it. Autism is a life-long condition and it is important to remember that people’s needs change as they go through life. Therefore, some people may need different types of support as they age.

Currently, individuals with autism may not find the services they specifically need or find them difficult to access. As such, they may miss out on achieving the things they wish to achieve in life. This is unfair and can lead to a poor quality of life and subsequent mental health problems.

Responsibility for delivering successful services for people with autism is widespread - there is not a singular agency that is responsible for providing services for people with autism. People with autism may be in contact with both specialist and mainstream services. This strategy will aim to make a link between specialist and mainstream services, to support coordinated responsibility and improve access to mainstream services.

We need an autism strategy to promote and develop change, in order to improve the quality of life for individuals with autism.

‘Never mind thinking outside the box, I just wish I could think inside the box’

(Person with autism)

We can do this through educating the population about autism and by making adjustments to our existing services to remove the barriers individuals with autism face and by further investing in and developing services across all sectors, who provide valuable services

An example of feedback in relation to ‘where we want to be’
There are a number of key national documents that are relevant to achieving the aims of our autism strategy. They impact upon the planning, commissioning and the delivery of a wide range of services. It is important for us to understand and incorporate these national initiatives, to ensure our local autism strategy reflects the strategic direction of the Scottish Government.

The Scottish Strategy for Autism, Scottish Government (2011)

‘Autism is a national priority’

Scottish Strategy for Autism, 2011

In 2011, the Scottish Strategy for Autism was published, declaring autism as a ‘national priority’, following a decade of autism specific initiatives (see Appendix A). The national strategy aims to harness these initiatives and address the entire autism spectrum and the whole lifespan of people living with ASD in Scotland, over the next ten years.

Considerable efforts have been made to improve diagnosis and assessment, to create consistent service standards, to match resources to need and to underpin this with appropriate research and training opportunities.

The vision of the Scottish Strategy for Autism is that “individuals on the autism spectrum are respected, accepted and valued by their communities and have confidence in services to treat them fairly so that they are able to have meaningful and satisfying lives.”

The Scottish Government also agreed the following values that underpin the National strategy:

- Dignity
- Privacy
- Choice
- Realising potential
- Safety
- Equality and diversity
The Scottish Government described 10 ways in which good services for people with autism should be developed:

1. A local strategy
2. Access to training and development
3. Easy access to useful and practical information about autism
4. An autism training plan
5. A process for data collection which improves the reporting of how many people with autism are receiving services and informs the planning of these services
6. A multi-agency care pathway for assessment, diagnosis and intervention
7. A way to get feedback to inform service improvement and encourage engagement
8. A multi-agency coordinated focus on meeting the needs of people with autism
9. Clear transitions at each important life stage
10. A self-evaluation framework to ensure best practice implementation and monitoring.

26 recommendations were listed in the Scottish strategy, which, in summary, said that:

- The Scottish Government would provide strategic leadership and create a strategic vision for the development of services and support for those with or affected by autism
- Resources would be effectively targeted with the aim of improving people’s lives
- People with autism and their families should be involved in decision making
- Cross agency working would be developed
- Adults should be able to get a diagnosis and support following a diagnosis
- People with autism should be supported to gain employment
The Ten Year Plan

To ensure the Scottish Government address their recommendations by the time the strategy concludes, they have set goals over 2, 5 and 10 year time periods. This is shown below and will guide the timeline of our local delivery plan.

<table>
<thead>
<tr>
<th>Foundations; 2 year goals</th>
<th>Whole Life Journey; 5 year goals</th>
<th>Holistic, Personalised Approaches; 10 year goals</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Access to mainstream services where these are appropriate to meet individual needs.</td>
<td>1. Integrated service provision across the lifespan to address the multi-dimensional aspects of autism.</td>
<td>1. Meaningful partnership between central and local government and the independent sector.</td>
</tr>
<tr>
<td>2. Access to services which understand and are able to meet the needs of people specifically related to their autism.</td>
<td>2. People with ASD have access to appropriate transition planning across the lifespan.</td>
<td>2. Creative and collaborative use of service budgets to meet individual need (irrespective of what entry route to the system is).</td>
</tr>
<tr>
<td>3. Removal of short term barriers such as unaddressed diagnoses and delayed intervention.</td>
<td>3. Consistent adoption of good practice guidance in key areas of education, health and social care across local authorities.</td>
<td>3. Access to appropriate assessment of needs throughout life.</td>
</tr>
<tr>
<td>4. Access to appropriate post-diagnostic support for individuals and families (particularly when there has been a late diagnosis).</td>
<td>4. Capacity and awareness-building in mainstream services to ensure people are met with recognition and understanding of autism.</td>
<td>4. Access to consistent levels of appropriate support across the lifespan including into older age.</td>
</tr>
<tr>
<td>5. Implementation of existing commissioning guidelines by local authorities, the NHS, and other relevant service providers. build skills and knowledge relating to alcohol and drugs</td>
<td></td>
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</tr>
</tbody>
</table>

Autism impacts on the whole life experience of people and their families; they need to be supported by a wide range of services such as social care, education, housing, employment and other community based services. A holistic, joined up approach is necessary.’

Scottish Strategy for Autism, 2011

Other national drivers that have influenced the development of our strategy and our action plan:

Social Care (Self-directed support) (Scotland) Act 2013

Self-directed support (SDS) is a term that describes the ways in which individuals and families can have informed choice about how their support is provided to them. It is most commonly used in the delivery of social care and support, but it can cover a much wider range of services.
SDS gives people control over an individual budget and allows them to choose how it is spent on support which meets their agreed health and social care outcomes.

SDS includes a number of options for getting support. The person’s individual budget can be:

- Taken as a Direct Payment (a cash payment);
- Allocated to a provider the individual chooses. The council or funding body holds the budget but the person is in charge of how it is spent (this is sometimes called an individual service fund);
- Or the individual can choose a council arranged service;
- Or the individual can choose a mix of these options for different types of support.

SDS allows people with autism and their carers to choose the support that is right for them and enables them to access a much greater range of supports to suit their needs and help them to achieve the outcomes that are important to them.

**Public Bodies (Joint Working) (Scotland) Bill (2014)**

This Bill seeks to achieve greater integration between health and social care services in order to improve outcomes for individuals and to improve the efficiency of services. There is no single definition of what constitutes integrated care, but the term is commonly used to refer to the joined up delivery of health and social care services. Integration is viewed as a way of tackling a number of problems such as unscheduled admissions to acute care, delayed discharges, budgetary battles between bodies, delays in accessing care and duplication of efforts. It is also seen as a way of ‘shifting the balance of care’, from the expensive acute sector, to care in less expensive community settings.

The concept of integration is not new to Scotland and the Bill is the latest in a line of attempts to achieve integrated care. Previous attempts have included the Joint Futures Agenda and the creation of Community Health Partnerships. The Bill proposes to require health boards and local authorities to create an integration plan for the local authority area.

This will be required for adult services, but other services may also be included. The integration plan will be required to detail which model of integration had been chosen and also sets out principles that should guide the creation of integration plans.

Integration proposals have the potential to greatly improve the way services are delivered for people with autism through a more joined up approach from diagnosis and treatment through to on-going support.


The Mental Health Strategy for Scotland indicates the priority placed on mental health by the Scottish Government and covers 14 high level outcomes. These include:

- People and communities protecting their mental well-being;
- People having a better understanding of their mental health;
- Care and treatment focuses on the whole person; and
- Professionals understanding the role of families and carers.

This single mental health strategy will encompass:

- Mental health improvement work;
- Mental illness prevention work; and
- Work to improve mental health services in general.

This will be a key influence in the coming years on the delivery of autism services and will help shape the services to include mental health improvement and the prevention of mental health problems.

The Scottish Government published ‘The Keys to Life’ in 2013, as a renewed and refreshed strategy for people with learning disabilities, following the publication of ‘the same as you?’ in 2000.

The Keys to Life sets out human-rights based principles, with aim of continuing a cultural shift towards meaningful change for people with learning disabilities, as well as continuing to improve quality of life for people with learning disabilities.

The strategy includes over 50 recommendations, embedded in 7 broad sections, including:

- Health;
- Independent living;
- Shifting the culture and keeping safe;
- Breaking the stereotypes;
- People with profound and multiple learning disabilities;
- Criminal justice;
- Complex care.

A local plan for people with a learning disability is currently in development in response to the National strategy.

Getting It Right for Every Child (GIRFEC), Scottish Government

The National Practice model is a dynamic and evolving process of assessment, analysis and review and a way to identify outcomes and solutions for individual children or young people. It allows practitioners to meet the Getting it right for every child core values and principles by being appropriate, proportionate and timely - (The Scottish Government)
GIRFEC in the context of the Scottish Borders Autism Strategy

The values and principles of GIRFEC will be recognised within the implementation of the priority areas identified in the Scottish Borders Council autism strategy. The National practice model will be used as a guide to implementing, monitoring and evaluating the actions from the strategy which relate to improving opportunities and experiences for children and young people living in the Scottish Borders. The local strategy also acknowledges the importance of practice which currently demonstrates how the GIRFEC framework is used within the region and the impact this has on outcomes for children and young people.


In 2001, the Public Health Institute of Scotland (PHIS) Autistic Spectrum Disorders Needs Assessment Report recommended that a SIGN guideline should be developed to improve the assessment and management of autism spectrum disorders (ASD) in Scotland.

The guideline applies to children and young people up to the age of 18, which may include the period of transition from childhood to adult services, focussing on assessment, diagnosis and clinical interventions for ASD. The guideline also considers joint working and consultation with children and young people, and with parents and carers. It also highlights how multidisciplinary and multiagency working can best address the needs of individuals with autism at all levels of service provision.


This guideline covers best practice guidance on the recognition and assessment of autism in adults. It also focusses on the care provided by primary, community, secondary, tertiary and other health and social care professionals who have direct contact with, and make decisions concerning the care of adults with autism.

Our strategy for the Borders will reflect and incorporate these priorities and national drivers; it is the responsibility of everyone involved to be mindful of these at the point of service planning and delivery.
CAN YOU SEE ME AUTISM AND ME
THE LOCAL PICTURE

Locally, there are policies in place which will inform the development and delivery of our autism strategy. Also, we can learn from previous autism specific projects that have been available locally, in order to help us identify what is working well, what needs improved and what is missing, for people with autism living in the Borders.

A description of these and their relevance are provided below; a link to each document can be found in Appendix B.

Local Policy

Scottish Borders Single Outcome Agreement (SOA), Scottish Borders Council (2013)

Scottish Borders Council and their community planning partners are responsible for working for the benefit of people living in the Borders. The outcomes of the council are published in the Single Outcome Agreement – a document agreed between the council and the Scottish Government.

‘By 2023, quality of life will have improved for those who are currently living within our most deprived communities, through a stronger economy and through targeted partnership action.’

The current SOA focuses on the key priority improvement areas for the Borders and highlights the areas where concerted partnership effort is required. These priority outcomes then have Performance Indicators. By monitoring things on a regular basis, it will be possible to see if the work being done by public sector partners is having a positive impact.


The Local Housing Strategy (LHS) provides the strategic direction to tackle housing need and demand and to inform the future investment in housing and related services across the Scottish Borders area. The LHS was developed through a consultation and community planning approach, setting out the key issues to be tackled over a five year period.

‘every person in the Scottish Borders has a home which is secure, affordable, in good condition, energy efficient, where they can live independently and be part of a vibrant community’

The Scottish Borders Local Housing Strategy 2012-2017
The LHS vision helps set the strategic tone for the document:

The LHS identifies four outcomes which Scottish Borders Council and its Community Planning partners are committed to delivering. These are closely aligned to a number of the Scottish Government’s national outcomes and there are direct linkages with the Scottish Borders Single Outcome Agreement. Outcome 4 is of particular relevance to our autism strategy: ‘More people with particular needs and/or requiring support are able to live independently in their own home’.

Children and Young People’s Services Plan 2012-2015, Scottish Borders Council (2012)

The Scottish Borders Children and Young People’s Planning Partnership (recently replaced with the Children & Young People’s Leadership Group) set out the key priorities for the group over three years, building on the outcomes from previous years’ plans.

The focus of the 2012-15 plan is centred on early intervention and the redesign of universal services to support this, through the continued implementation of GIRFEC. The priority areas identified include:

- Early years
- Promoting children’s rights
- Keeping children safe
- Looked After and Accommodated Children and Young People
- Parenting
- Parental involvement
- Improved attainment and achievement for all our children and young people
- Improved health and wellbeing for children and young people
- Transitions 16+
- Workforce planning and development

The subgroups of the Children & Young People’s Leadership Group have responsibility for delivering on these key priority areas.


The Scottish Borders Youth Voice engaged with Borders young people to collect key messages about the provision of health services in the area and to identify their expectations for health practitioners. The strategy describes how NHS Borders plans to further improve the health and wellbeing of children and young people up to the age of 18 years old.

Five outcomes are identified in the health strategy:

- Have the best possible start in life and improvement in their wellbeing
- Have access to high quality person and family centred health care at the right time and in the right place
- Receive care and support that is targeted for those who are vulnerable and at risk of poor health outcomes, including mental health
- Be involved in decisions and planning that affect their health and when appropriate include their families too
- Have an improved experience for their transition into adult health services

Local policy and strategy development needs to ensure that services for people with autism are inclusive and meet all equality and diversity strands. Links between the autism strategy and local policy priorities should be developed and strengthened as part of improving the outcomes for people with autism living in the Scottish Borders.
Previous local Autism specific projects


From 2005 to 2006, the Scottish Executive (Scottish Government) funded the Adult Autism Spectrum Disorder (ASD) Co-ordinator post. The remit of the co-ordinator was firstly to be a ‘change agent’ in the lives of adults with ASD, their families, friends, carers and the services that support them in the Scottish Borders. Second, the co-ordinator was responsible to help plan, in partnership with NHS Borders and Scottish Borders Council, services for adults with ASD and their families.

There were 8 key areas to action:

- Driving forward the agenda for adults with ASD;
- Audit the numbers of adults with ASD in the Scottish Borders;
- Promote the development of an ASD database;
- Audit services and provision available to adults with ASD;
- Create ASD information base;
- Promote the use of a multi-agency care pathway for adults with ASD and the use of Person Centred Planning;
- Raise awareness of ASD in the Scottish Borders;
- Promote the review of adults with ASD who are placed outwith the Scottish Borders.

The Autism Co-ordinator project was the beginning of a number of positive initiatives and was key in raising the profile of adults with autism living in the Scottish Borders.


As part of the National strategy launch, the Scottish Government announced £13.4 million of additional investment to be distributed over 4 years, some of which would be dedicated to mapping out autism services and improving coordination of these services.

Coinciding with a one-off investment of £35k for each local authority in Scotland to develop their own local Autism Action Plans, the aim of the Autism Mapping Project (‘the project’) was to:

- consult with people with autism, their families and carers, service providers and local agencies;
- map out existing autism service provision in all local areas in order to build up a local and national picture;
- identify priority areas for action that reflected local need;
- work collaboratively with local partnerships, councils, NHS, criminal justice, third sector organisations and other relevant public bodies;
- provide local authority with a ‘Service Map’ of their area, to inform their Autism Action Plans.

A national mapping report was developed which provides a ‘snapshot’ of autism services across Scotland, setting out the key issues identified by people with autism and their carers, and provides an overview of how services are meeting their needs or where there may be gaps in services.

The local autism service map was compiled using evidence collected in the Scottish Borders. We asked people with autism, parents & carers and professionals for their views on what is working well, what needs improved and for any gaps in service provision in the local area.
The data collected from the workshops and the questionnaires was analysed to create a number of key themes across all key stakeholder groups.

The key themes from each stakeholder group [people with autism, parents & carers, and professionals] were compared and the priorities that matched became our initial priorities, for further engagement.
Collecting data about autism can be used to promote early identification, plan for training and service needs, guide research and inform policy, so that people with autism and their families get the help they need. It is difficult to gather data accurately, due to the range of information recording systems used between services and professionals. As a result, there can be instances where data is absent, or is counted more than once. It is important to also note that a number of people with autism exist in the population whom have never been formally diagnosed, and as such, will not be represented in the data recorded.

**National estimated prevalence**

The prevalence estimates in the Public Health Institute of Scotland (PHIS) ASD Needs Assessment Report (2001) suggested 60 in every 10,000 people have autism. However, the Scottish Strategy for Autism (2011) reports that more recent studies estimate the prevalence of autism in children and adults in Scotland to be around 1 in 100.

Autism is diagnosed more commonly in males than females, at a ratio of approximately 4:1, although this varies across the spectrum.

**Local estimated prevalence and recorded data**

According to Scotland’s 2011 Census, the population of the Scottish Borders is 113,870. Using the estimated prevalence of 90 per 10,000, as detailed in the Scottish Strategy for Autism, there would be an estimated 1,025 people with autism living in the Scottish Borders.

**Adults**

The Scottish Consortium for Learning Disabilities (SCLD) reports in the 2013 release of Learning Disability Statistics Scotland (eSAY) report that there are 3,655 adults in Scotland with autism spectrum disorder who are known to local authorities; most of these adults have a learning disability.

SCLD reports that in the Scottish Borders, there are 107 adults identified with an autism diagnosis. This includes: 98 people with a diagnosis of ‘Classic Autism’; and 9 people with a diagnosis of ‘Asperger’s Syndrome’.

**Children and Young people**

The Additional Support for Learning and Young Carers Report for Parliament (2013) reports that in the Scottish Borders there are 1,757 children and young people in education with additional support needs. The report highlights that 76 of these individuals have autism in the Scottish Borders.
**People with autism known to statutory services**

We asked Scottish Borders Council and NHS Borders to collate the number of people with a diagnosis of autism (including Asperger Syndrome) known to their services.

**SBC Social Work data**

The table below shows the number of people in contact with social work services. Please note:

- The people represented below may or may not have a formal, clinical diagnosis of autism or Asperger Syndrome;
- Some people with autism who have had contact with social work services may not appear in the table, due to recording methods;
- Some people with a diagnosis of autism or Asperger Syndrome are recorded in the NHS data, but have not been in contact with SBC Social Work, so will not be represented in the table below;
- These figures represent a snapshot of data currently held

<table>
<thead>
<tr>
<th></th>
<th>Autism</th>
<th></th>
<th>Asperger Syndrome</th>
<th></th>
<th>Grand Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Female</td>
<td>Male</td>
<td>Total</td>
<td>Female</td>
<td>Male</td>
</tr>
<tr>
<td>Children under 16</td>
<td>1</td>
<td>27</td>
<td>28</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Adult 16+</td>
<td>21</td>
<td>68</td>
<td>87</td>
<td>2</td>
<td>9</td>
</tr>
<tr>
<td>Grand Total</td>
<td>22</td>
<td>95</td>
<td>117</td>
<td>2</td>
<td>11</td>
</tr>
</tbody>
</table>

**NHS Borders data**

**Children**

The multi-agency team situated within the Child and Adolescent Mental Health Service (CAMHS) provides assessment of children and young people with possible autism up to the age of 18 years.

From 2009 to 2014, there were 114 children referred to this service, and 60 were diagnosed with autism.

**Adults**

The NHS Borders data supplied gives the number of people with autism who have had an inpatient episode, or have been seen by the Mental Health Service, or the Joint Learning Disability Service. Data on wider prevalence is not currently available.

<table>
<thead>
<tr>
<th></th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Joint LD Service</td>
<td>29</td>
</tr>
<tr>
<td>Annual Mental Health Teams</td>
<td>40</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>69</td>
</tr>
</tbody>
</table>

Wherever possible, we have tried to avoid duplication of numbers within the data presented; however, due to the recording methods used, a degree of duplication may exist.

For information regarding spend across agencies, please refer to Appendix B.
Our autism strategy represents extensive engagement with key stakeholder groups. Their feedback has directly shaped the content of our strategy and has signalled the changes required for better outcomes for people with autism and their families.

Listed below are the steps taken to ensure our strategy was successfully co-developed with those it would affect.

**Step 1: Identifying our priorities**

A strategic group have been meeting since June 2012 to identify areas of need and develop action points to address them. Members of the strategic group include representatives from NHS Borders, Scottish Borders Council, education (including further education), learning disability services, the voluntary sector, a carer organisation and parents of people with autism.

We used the local Borders Service Map from the Autism Mapping Project (for a full description, see ‘Local Autism Specific Projects’ above) to help us identify our initial priorities for further development.

**Step 2: Asking people what they thought of our initial priorities**

Over a 6 month period, we engaged with a range of key stakeholders, including senior managers, practitioners, clinicians, parents and carers, and people with autism. Using a variety of methods, including meetings, interviews, questionnaires and interviews, we asked people if they agreed with our initial priorities and for their opinions and personal experiences related to them. The table below shows the record of engagement.

<table>
<thead>
<tr>
<th>Group meetings</th>
<th>Individual interviews</th>
<th>Workshops</th>
<th>Questionnaires</th>
</tr>
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<td>2</td>
<td>6</td>
<td>2 adult groups&lt;br&gt;2 children and young people groups</td>
<td>170 responses&lt;br&gt;10 easy read responses</td>
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</table>

**Step 3: Incorporating people’s views into our strategy**

The statements and content detailed under each priority area in the following pages of the strategy have been drawn from all stages of the engagement process, including our questionnaire, workshops, meetings and interviews.

The feedback was summarised and the main themes are included as part of the narrative running throughout the strategy.
Step 4: Final draft version consultation

Once the final draft version of the strategy was finished, we launched a 12 week consultation period beginning on World Autism Awareness Day on 2nd of April 2014, to find out what people thought of our strategy. We attended group meetings, held an individual interview and made consultation versions of the strategy available for feedback. The table below shows our record of consultation.

<table>
<thead>
<tr>
<th>Group meetings</th>
<th>Individual interviews</th>
<th>Feedback forms</th>
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<tbody>
<tr>
<td>13</td>
<td>1</td>
<td>8 responses</td>
</tr>
<tr>
<td></td>
<td></td>
<td>4 Easy Read responses</td>
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</table>

The feedback was collected and analysed, with any necessary changes made for the final version of our strategy.
WHAT IS WORKING WELL?

In order for our strategy to be successful and improve the lives of people with autism, we must learn from, and continue to build on, the vast amount of good work that already exists here in the Borders.

Engagement with people with autism, their carers and families and professionals, during both the mapping project and through the development of the strategy, highlighted a number of examples of effective autism practice. Below are two examples selected to reflect the developments of effective practice in the early years.

Effective practice example 1: Co-ordinated Support in Education

A number of positive experiences were reported within primary education. An Individualised Education Programme/plan (IEP) is key in coordinating various strands of support: input from Speech and Language therapy and Occupational therapy have a significant impact on the educational experience of the child.

Additional adjustments made by a teacher whom has attended autism awareness training were also highly beneficial. Positive approaches include ear defenders, having a quiet place to go, and supporting the child as an individual, rather than a diagnosis. Good practice was also evident in the communication between professionals around the IEP.

Effective practice example 2: Transition Planning for Child in Primary Education

Spectrum Support, the education outreach team ensure that a dedicated transitions teacher is allocated to each P6 pupil with autism at the very start of the primary/secondary transfer process. The teacher gets to know the child in their familiar primary setting before introducing them to their local secondary school where they will continue to support them throughout their secondary career.

The support includes: supported visits; developing pupil profiles; transition booklets; working with parents; supporting independent travel to their new school; and training of school staff in the receiving school prior to them starting. This ensures children and their parents/carers are well-informed about the new school, have a known face to continue support in the new setting and school staff have been prepared for their new pupils.
We will work together towards achieving our vision, over the next 10 years.

We have engaged with key stakeholder groups and asked them what they think is working well, what needs to improve and what gaps exist for people with autism living in the Borders. This information has helped us identify the foundations of good practice to build upon, and other areas of life that we will aim to develop. The themes for further development were cross referenced between each stakeholder group and we identified 7 specific, consistently referenced priorities.

The following sections will detail what people have said about each theme, and the call out boxes show direct quotes from questionnaire respondents. The resulting actions for change can be found in the corresponding thematic section in the action plan:

1. Autism awareness and training – Improving public and professional awareness and understanding of autism;

2. Diagnosis – Good-quality, early diagnosis and intervention for both children and adults;

3. Getting the right services at the right time, for adults with autism and no learning disability – Addressing the eligibility criteria and improving access to appropriate support;

4. Purposeful occupational activities – Finding the right opportunities to pursue individual interests and employment;

5. Social support and opportunities – Opportunities to develop social skills and access a range of social opportunities;

6. Improving access and provision of housing – Addressing the barriers to finding, securing and maintaining individual housing needs;

7. Ensuring inclusion for people with autism and their families – People will have their voices heard and acknowledged by professionals.
1. Autism awareness and training

*Improving public and professional awareness and understanding of autism*

“This is very important for a number of reasons; early intervention, management of the condition, and understanding how, very often, people on the spectrum see and understand the world in a very different way from someone who is not on the spectrum.”

Questionnaire respondent

**Feedback Received Summarised**

Over 90% of the respondents to our questionnaire agreed that autism awareness and training should be a priority for further development in the Borders. People stressed the importance of increasing autism awareness and understanding in order to make a significant difference to the life of people with autism.

Improved awareness and understanding of autism can improve access to key, mainstream services. There is evidence which shows that basic awareness training on autism can significantly improve people’s ability to communicate with people with autism. Developing a better understanding of autism will also address commonly held misconceptions in society, which can be potentially harmful.

Autism is a complex condition and, for some people, it may not seem obvious that they have a diagnosis – some people refer to autism as an ‘invisible condition’. People commented that autism awareness initiatives should specifically include information about Asperger Syndrome and the difficulties somebody may experience in their daily life.

Through our engagement, people highlighted the need for society in general, and both mainstream and specialist services to develop a better understanding of autism, for people with autism to lead fulfilling lives. There is a lot of good, individual work that exists already to increase autism awareness, however people are not always aware of it and it can often be inconsistent. People with autism said there needs to be a shared understanding of autism across both NHS services and Scottish Borders Council.

“There is a lot of good individual work, however there needs to be a cohesive approach used”

The graph below shows services people specifically named that they feel need further autism training. Some of the responses were from service providers themselves, demonstrating the people working in the field are keen to develop their own autism knowledge.

The responses were grouped into relevant agencies – for the full list of these services, please see Appendix C. The table shows that statutory services (NHS Borders, Education and Other council services) are among the top three agencies in need of further autism training.
Where we want to be

The NHS Education for Scotland Autism Training Framework, ‘Optimising Outcomes’, provides a scaffold for all staff working with people with autism, their families and carers. The framework describes three key areas relevant to Health and Social Care settings. Within these areas, four levels of knowledge and skills exist, dependent on the varying degrees and frequency of contact with people with autism and their families, ranging from the primary level of autism knowledge towards the specialised level of autism knowledge:

1. Autism informed Practice Level
2. Autism skilled Practice Level
3. Autism enhanced Practice Level
4. Expertise in autism Practice Level

Staff can identify their required level of autism knowledge and skill using the Autism Training Framework within their organisation; appropriate training must then be identified and accessed. This tool should be utilised in the Borders, to allow the identification of training needs for the individual or service. People with autism could be involved in the development and the delivery of training.

We want the general population to recognise autism. This begins with a broad approach to developing awareness in society, represented in the first tier of the triangle figure below. Some people suggested using creative and innovative methods of increasing autism awareness locally; this could include plays or drama shows.

In order to ensure services meet the needs of people with autism, service providers need to be targeted with an in-depth level of autism training. This would apply to some of the services shown in the graph above and is represented as the middle tier of the triangle figure. The Training Framework described above will be instrumental in achieving the appropriate levels of autism knowledge and skills required by this group of professionals.

The top tier of the triangle represents the highest level of training for specialist professionals working in the autism field. In order for services to be equipped with the most up to date information, specialist practitioners need to continue to receive this level of training. Again, the Training Framework provides specific criteria for this group of professionals and will be important in determining the training requirements.

This visual model of training shows the various levels of knowledge and skills that should be available locally; ranging from the broad autism awareness and basic level training for mainstream services and the community, to high-level training intended for specialists.
2. Diagnosis

**Good-quality, early diagnosis and intervention for both children and adults**

*The earlier autism can be diagnosed and assessed, the better the quality of life for the individual*

**Feedback Received Summarised**

Over 95% of questionnaire respondents support the diagnostic assessment of autism as an area for future development.

Going through an assessment for autism and receiving (or not receiving) a diagnosis can be both a positive personal experience and a stressful, emotive experience. We know that having a diagnosis of autism can be the first step towards self-understanding and can be the key to accessing the right support.

Early assessment and intervention are reported as significantly improving outcomes for people with autism. However, feedback showed that some people had difficulty being referred for an autism assessment and others had to wait a long time for the assessment. In some cases, people were diagnosed much later in life after going through various other routes in mental health services.

**Pre-diagnostic awareness and support**

Feedback indicated some people are not sure of any existing pathways or routes to and through the diagnostic process and do not know where to look to find this information. We are aware that a pathway exists for children, however not everyone is aware of this. Therefore, people are often not sure how to start going about obtaining an assessment for autism. This often led to people carving out their own path to an autism assessment, which prolonged the process greatly. Some feedback showed that for adults without a learning disability, the route to diagnosis is especially not clear. Although the process begins with an appointment with the GP, the next steps are still not well known.

*Clear guidelines should be available as to how to access an assessment, what it will involve and how long the process takes*
People thought that regular autism training for frontline staff across a number of agencies (including health visitors, community based services, nursery/primary/secondary teachers, and adult service practitioners) and the public in general, would help people recognise autism and direct people to the next action, towards an autism assessment. It was thought that this would allow for a quicker referral to the relevant diagnostic service.

The Assessment Process

Going through an assessment for autism can be very stressful for the individual and their family. It was reported that there is a lack of support during the assessment process. People said that information about the assessment and what to expect would be helpful in reducing stress and anxiety.

We have been told that for adults without a learning disability, the assessment process is very difficult and there is a need for a local service, dedicated to assessing this client group.

People also talked about the length of time the assessment takes to complete; some people thought it took too long and wanted to know the outcome quicker than is currently happening. Feedback from clinicians, however, indicates that the length of assessment is dependent on the complexity of the assessment and the number of professionals involved, due to the multi-disciplined nature of the assessment.

Post-Diagnostic support

People highlighted both short term and long term intervention as important. Key to both these timescales is information; people want information to be made available on various mediums, including websites, mobile phone apps, books and a list of key contacts and services in a directory. Access to appropriate services for the short term and long term planning was highlighted as important. Furthermore, people were not sure of any existing pathways which map out an individual’s route from diagnosis to existing services, particularly in adult services. It was also reported that families should receive information for where they can find support, to ensure a holistic approach is taken to support.

Feedback included the need for regular reviews to occur, following a diagnosis, in order to plan out key life stages.

Voluntary organisations were reported as being supportive and helpful following diagnosis; people are keen that the assessment service develop stronger links with existing support services, in the voluntary sector.

Where we want to be

We want people to have easy access to a single point of information about the diagnostic process, including pathways, so they are clear about who to contact, what the assessment will involve and what will happen after the assessment; irrespective of whether or not they have an existing mental health condition or learning disability. This should be available in a variety of formats and should link together NHS Borders and Scottish Borders Council information. It is hoped that with clearer pathways, transitions between services and professionals is more effective and smoother.

We want the assessment process to be completed to a high and consistent standard, in line with current clinical guidance. People should be informed about what to expect and be aware of the estimated time periods to complete the assessment.

We want people to have equal access to appropriate support following a diagnosis of autism, with information about the condition and about available services.
3. Getting the right services at the right time, for adults with autism and no learning disability

*Addressing the eligibility criteria and improving access to appropriate support*

"I feel there is a lack of support and services for people with autism and no learning disability. People assume if you are of high intelligence you do not require any support."

**Feedback Received Summarised**

Over 90% of respondents agreed that getting the right services at the right time, for adults with autism and no learning disability should be a priority for further development. This client group are frequently known as having Asperger Syndrome or High-Functioning Autism.

The needs of this group of people can be intricate and complex; their needs can be difficult to identify through typical assessment and can be dependent on environmental factors.

People reported a gap in services for this client group; often, people with autism and no learning disability fall in between mental health services and learning disability services. Some people told us the needs of this client group are not fully understood and are not incorporated well in social work assessment tools as a result, which means they often miss out on the support they need.

We understand that people are not aware of existing services, available locally or how to access them; feedback indicated a lack of information also contributed to the feeling of a lack of services in the Borders.

People described a range of services that were required for this client group to work towards independent living, including: post-diagnostic support; employment support; benefits advice; befriending; social activities; housing; and further education support. The benefits of having an expert level of autism knowledge in these services was apparent through our engagement.

The geography of the Borders was raised as a potential barrier for finding a central location for services, which could cater for a number of rural towns.

**Where we want to be**

The needs of people with autism and no learning disability are sometimes referred to as ‘hidden’ or ‘invisible’. This client group are more likely to be diagnosed later in life, or will remain undiagnosed than those with autism and a learning disability. Therefore, this group is often disadvantaged when having their needs recognised and met by services.

We want understanding of autism and no learning disability (commonly referred to as Asperger Syndrome or High Functioning Autism) to improve greatly, so that the needs of this client group are accurately assessed and supported. We want people to know where to go to find information, that is accessible and up-to-date, and for people to know how to access services that are right for the individual. We also want professionals to know where to refer individuals to, at appropriate times, to ease transitions between services and professionals.
4. Purposeful occupational

Finding the right opportunities to pursue individual interests and employment

*Autism ceases to be a label when you have a purposeful occupational activity. You are what you do.*

Questionnaire respondent

Feedback Received Summarised

Being able to find and pursue individual interests and employment was regarded as ‘critical’ to young people’s lives, and over 90% of questionnaire respondents agreed this should be a priority for future development in the Borders.

Leaving school and knowing the options you have is important. Being able to pursue your interests as an adult, in a meaningful way, either through further education/training or employment, is also vital.

People have told us that they sometimes do not know the options they have upon leaving school and this can be a source of stress and anxiety for both the individual with autism and their families. Transitions in life are referenced frequently as being key to securing good outcomes and minimising stress and anxiety.

There are barriers which can prevent people with autism from accessing further education, training and occupational opportunities that were reported through our engagement. Feedback showed that a lack of understanding of autism from employers is thought to be the most significant barrier to obtaining and maintaining employment.

Voluntary work was cited frequently as a good step towards gaining paid employment and a way to actively engage in a personal interest.

It was recognised that in order to be successful in training or further education, some people require additional support, to help manage anxiety and social issues that may arise during the course or work placement. Getting the right support from employment staff who are knowledgeable about autism, to help find and apply for training and jobs, was also raised as a priority. Autism specific courses were also raised as a method of ensuring access to training was made easier.

It was also reported that people with autism would benefit from having support available while in employment, to help maintain their role.
Where we want to be

Having a purposeful occupation and activities can provide people with a sense of meaning, enjoyment and social opportunities. In a broader sense, purposeful occupation provides society with economic contributors and this can be valuable to the local community.

We want life-long planning, beginning in education, for people to map out options based on their personal strengths, skills and interests to improve transitions and outcomes. We also want local employers to develop a better understanding of autism, to improve equal access to opportunities for employment.

5. Social support and opportunities

Opportunities to develop social skills and accessing a range of social opportunities

“Great idea. Of course I crave social activities, like most folk, but do find it difficult, often entering into conflict, I am often unaware of why these conflicts develop, and I seem to have an amazing ability to upset certain people without understanding how” – Questionnaire respondent

Feedback Received Summarised

Developing more social opportunities and support in the Borders is a priority for the future, with over 90% of questionnaire respondents in agreement.

Social interaction can be a very challenging and stressful experience for people with autism; however this does not mean that all people with autism do not want to mix with other people. The feedback demonstrated that having the choice to socialise was important, and that ‘neurotypical’ values should not be imposed on this client group.

For some people with autism, having a range of social opportunities is important; ranging from developing community based activities which are tailored to the individual, to more supported means of socialisation. Some people advocated the development of autism specific social services, as a way of interacting with people with the same condition.

“It was also thought that the public have a responsibility to become more accepting and understanding of autism, through awareness raising, so that existing community based activities become more accessible for people with autism.”

“We could establish an effective network of professionals already providing employment support, so that repetition is avoided and perhaps some specialisms could be developed in terms of the type of support on offer and at what point in the employment pathway that support is given.”

“Having a range of options and choices is important. Focus on service user involvement, supported as necessary.”

“The social side can be the hardest bit for those with autism. It is important to find social places that they feel comfortable and to have support workers to help make sure they are in the community more to help with social situations”

“The general public need to adapt to the needs of others, particularly people with autism, rather than force those with autism to develop social skills with which they have less of a connection to the neuro-typical psychology.”
There is evidence to show that many people with autism feel that with appropriate social support, quality of life would drastically improve and mental health issues would reduce. Some people said having the opportunity to share their favourite activity with others would help manage their anxiety and provide them with a chance to meet other people.

The feedback shows that some people with autism need specialist social support, available locally, to learn aspects of socialisation that would allow them to access a greater variety of existing opportunities, for children, teenagers and adults. Transitioning from social activities in children’s services to over-16 opportunities was also thought to be key.

Peer mentoring and ‘buddying’ or befriending schemes in the voluntary sector received very positive feedback; the one to one nature of the scheme was recognised as essential in working towards broader inclusion and building confidence.

Where we want to be

We want people with autism to have access to a range of social opportunities, which are genuine, varied, and tailored to meet the need and expectation of the individual. Having choice in the activities you do is vital. The feedback showed that in order to do this, partner agencies need to work together, to make transitioning between services better and to overcome potential funding problems people expressed.

We want people with autism to have more opportunities to learn social skills, in a supported environment, should this suit the individual.

We also recognise that social opportunities need to be available locally; as the geography of the Borders can present numerous challenges to travel, resulting in isolation.

We also want to address the existing barriers to wider community based activities that people with autism want to access.

6. Improving access to and provision of housing

Addressing the barriers to finding, securing and maintaining individual housing needs

“the light at the end of the tunnel is knowing that the person you care for is in appropriate housing for their needs”

Questionnaire respondent

Feedback Received Summarised

Moving out of the family home and having a flat or house of your own is seen as a fundamental step towards living independently. For people with autism, this can be challenging, for a number of reasons. Our questionnaire shows that over 90% of respondents agree that improving access and developing the provision of housing in the Borders should be a priority for future development.

Some people told us that they felt there is a lack of appropriate housing options for people with autism in the Borders, which meant that individuals were staying at home with their parents for a longer period of time than is desired.
Some people expressed a need for increased housing stock, available locally, built with the needs of people with autism in mind at the point of planning.

The feedback also reflected that a range of housing options are required, to meet the various levels of need associated with autism. Some people referred to the existing supported accommodation as working well, however others reported a need for more specialist provision to be made available. On the other hand, feedback also highlighted a need for more independent living opportunities, with support to apply for and manage tenancies.

The process for applying for social housing was raised by a significant number of people through our engagement. People felt that the needs associated with autism were not always taken into account when finding or applying for housing. It was thought that the vulnerability of some people with autism should be acknowledged in this process, which should influence the potential areas selected for living.

Where we want to be

We want people with autism to have equal access to a range of housing options, which are carefully selected, to meet the unique needs of the individual. We want people to know where to access information about housing and who to contact for advice. Again, it is vital for all parties involved to be clear about available housing options to ensure transitions are as simple as possible. Transitions between housing circumstances need to be understood and supported effectively.

We also want to work with our partners, to look to the future, in developing more appropriate housing which has been planned with autism in mind.

7. Ensuring inclusion for people with autism and their families

*People will have their voices heard and acknowledged by professionals*

"I think it is a great idea. The people who require, and will be using the services are the best people to be involved with these decisions. They are the only ones who can give personal opinions, feedback and advice on current and future services."

It is essential that the views of people with autism and their families are heard and acted upon. Over 95% of respondents to our questionnaire agree that further developing inclusion should be an area of development in the Borders, as part of our autism strategy.

People highlighted good practice that exists in education, through the Education Act Scotland, which ensures equality and inclusion. The ‘Meeting Around the Child’ meetings and the Children’s panel are reported examples of this good practice.

The feedback clearly shows that people want to be heard; people told us that listening to parents, carers and people with autism is key to developing tailored services and meeting the needs of the individual. Another key point reported is the need to transfer information collected from stakeholders into action, to make real changes.
People suggested a range of methods for how they could best express their views and opinions. The graph below shows the methods suggested and the number of people that suggested the method.

From the data, it is clear that online methods are considered a popular, accessible way of sharing opinions and views. It was also reported that everybody has access to the internet or wants to access the internet and as such, other means of communication should be used as well.

The feedback also stated that effort should be made to find the most suitable method of communication, which might include using writing or pictorial based communication methods, for the individual.

“There is no ‘best’ way (to communicate) as people on the spectrum don’t all respond in the same way – needs an individualised, person centred plan”

Where we want to be

We want people with autism, and their parents and carers, to have equal opportunities to express their opinions and views to knowledgeable professionals, to help manage and plan existing and future services. We want a range of communication methods to be available, to empower people with autism to have choice, and ensure their voices are heard and taken into account in the future.
Autism and me
I am more happy mummy
The development of our autism strategy has given us a real sense of what people think the future should look like for people with autism and their families living in the Borders. The priorities section listed above reflect the feedback we received, including what people felt was working well at the moment, and where we want to be in the future. Our vision will serve to represent these principles and values moving forward with our progress.

The next steps will involve a great deal of work, from a range of partners. Our delivery plan will help guide this work over the next ten years and aims to provide potential actions collected through our engagement and consultation. The delivery plan is a live, dynamic document and will be updated accordingly as progress develops.

We would like to thank everybody whom made a contribution to the development of our strategy. Their feedback has proven to be invaluable and has resulted in an organic document which will serve to improve outcomes for people with autism and their families living in the Borders.
APPENDIX A

List of previous autism initiatives, cited in the Scottish Strategy for Autism


SIGN publication No 98. Assessment, diagnosis and clinical interventions for children and young people with autism spectrum disorders
http://www.sign.ac.uk/guidelines/fulltext/98/index.html

Commissioners of health and social care services for people on the autism spectrum, Policy and Practice Guidance, (2008)
http://scotland.gov.uk/Publications/2008/03/27085247/0


http://www.scotland.gov.uk/Publications/2008/03/27092411/0

A Guide to Getting it right for every child (GIRFEC), (2008)
http://www.scotland.gov.uk/Topics/People/Young-People/gettingitright
**APPENDIX B**

**Breakdown of current estimated spend on people with autism:**

It has been extremely difficult to calculate the current spend on support and services for people with autism. All figures below are indicative only and apply to the current financial year only and may not necessarily be representative of actual spend.

**Joint Learning Disability Service:**

The Joint Learning Disability Service calculates that current spend per annum on people with a learning disability and also autism is **£3,202,503**. This comprises of individual packages of care for **59 clients** which equates to an average cost of **£52,690** per person. The types of support provided within this sum are community support, housing support, personal care, day care, residential as well as other services.

**Access to Number 6 One Stop Shop:**

The cost is currently **£15,000** per annum to provide access to people with Asperger Syndrome and High Functioning Autism from the Borders.

**Staffing Costs:**

It has proved difficult to calculate costs relating to staffing time devoted to working with people with autism. An indicative or proxy figure of **£24,000** per annum has been given as the estimated costs of time devoted by three NHS posts involved in diagnosis and assessment.

It is impossible to calculate what the indicative costs will be, linked to all the time spent by a wide range of NHS and SBC staff in their working with people with autism.
Full list of services specifically named by questionnaire respondents as in need of further autism training:

All staff/front line employees
Medical
Community/public
Podiatrists
Employment support
GP/Doctors
Hospital
Legal
Health
Nurseries
Supermarkets
Cinema
Restaurants
Community services
Out of school clubs
Housing officers
Modern technology
Befrienders
Advocacy services
Skills Development Scotland
Psychiatrists

Teachers
Police
Dentists
School
Call centres
Social Work
Education
Leisure/recreational
Charitable organisations/3rd sector
Local businesses/private sector
Cafes
Libraries
Mental Health Services
Childcare
Pharmacy workers
Carers
College/University
Health visitors
Citizens Advice
Psychiatric nurses
Job Centre Plus Staff
REFERENCES

http://www.scotborders.gov.uk/info/828/activities_and_support_for_young_people/141/strategies_for_children_and_young_people

Getting it Right for Every Child (GIRFEC), Scottish Government
http://www.scotland.gov.uk/Topics/People/Young-People/gettingitright

http://www.scotland.gov.uk/publications/2012/08/9714


http://www.nice.org.uk/guidance/cg142

Public Bodies (Joint Working) (Scotland) Bill (2014)


http://www.scotborders.gov.uk/downloads/file/6949/national_autism_services_mapping_project_scottish_borders_service_map


http://www.scotborders.gov.uk/info/691/council_performance/1351/single_outcome_agreement
Scottish National Census, Scottish Government, [2012]
http://www.scotlandscensus.gov.uk/

Social Care (Self-directed Support) (Scotland) Act, [2013]
http://www.legislation.gov.uk/asp/2013/1/contents/enacted


The Keys to Life - Improving Quality of Life for People with Learning Disabilities, [2013]
http://www.scotland.gov.uk/Publications/2013/06/1123


The Scottish Strategy for Autism, Scottish Government, [2011]
http://www.scotland.gov.uk/Publications/2011/11/01120340/0
Agency – A team or group of individuals delivering a specific type of service within the council or National Health Service.

Anxiety – Feeling particularly worried or concerned about something with an uncertain outcome.

Asperger Syndrome – A form of autism, with no cognitive or speech and language development delay, and an Intellectual quotient (IQ) of at least 70.

Atypical autism – Used when the person’s behaviour pattern fits most but not all the criteria for typical autism.

Co-existing Conditions – Having more than one health condition.

Cognitive style – The way that somebody thinks; perceiving, interpreting and understanding information.

Consultation – Using different methods of asking questions to collect general feedback from people.

Engagement – Using different methods of asking questions to collect feedback from people about specific topics.

Flexibility of Thought – The cognitive ability to adjust ideas and expectations based on new information.

High Functioning Autism – A form of autism, with significant delay in the development of early speech and language skills, and an IQ of at least 70.

Housing stock – The volume/quantity of available housing.

Intellectual quotient (IQ) - A score derived from one of several standardized tests designed to assess/measure human intelligence.

Language processing - The cognitive processing of spoken or written language. This ranges from the construction of spoken or written messages to the abstract meaning of language.

Learning disability - Significantly reduced ability to understand new or complex information, to learn new skills and reduced ability to cope independently which starts before adulthood with lasting effects on development.

Mainstream services/Universal services – Services designed for the entire population.
**Multi-agency** – A group of professionals representing a variety of service teams.

**Neuro-typical** – A term often used in the autistic community and field, as a description for people who are not on the autism spectrum.

**Outcome** – A meaningful change experienced by an individual.

**Pathways** – A specific and agreed order of steps involving different professionals and services to attain an outcome.

**Pervasive Developmental Disorder Not Otherwise Specified (PDD-NOS)** – Used when the person’s behaviour pattern fits most but not all the criteria for typical autism.

**Prevalence** – The reported proportion of a population that share a common condition or set of characteristics.

**Proprioception** – Sensing your own body in relation to the physical space; having body awareness.

**Priorities** – Areas in need of development and attention.

**Specialist services** – Services designed for a specific section of the population.

**Third sector organisations** – Charitable organisations providing support services.

**Transition** – Moving from one experience to another.

**Vision** – A shared belief of what we should strive to achieve.

**Vulnerable** – Being at risk of a negative experience.

**Wellbeing** – The state of being comfortable, happy, or healthy.