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Since I began my role as National Clinical Director for Diabetes, I have been repeating a message to the diabetes community: *Every person with diabetes deserves the highest standards of care, no matter where, when or by whom that care is delivered. Healthcare professionals delivering such care should be trained in diabetes, and recognise the boundaries of their knowledge. They should be given opportunities to extend those boundaries, and to update. Professionals and people with diabetes must have ready access to specialist advice.*

Of course this message applies not just to people with diabetes, but with any health condition. The elements needed to ensure equal access to such care are beginning to extend in more and more NHS communities.

First of all, it is essential that the services provided are comprehensive and seamless, looking at the whole needs of the person and working across financial and organisational boundaries, e.g. across primary and secondary care. As an example, a working party comprising representatives of the Royal College of Physicians of London, the Royal College of General Practitioners and the Royal College of Paediatrics and Child Health, supported by the NHS Alliance, has recently published the excellent *Teams without Walls*, which brings with it the potential for seamless multidisciplinary care for people with long-term conditions.

But there is also another shift underway – in the relationship of people with long-term conditions to the service itself. In this era of personalisation and choice, people with long-term conditions are more and more being recognised as experts in their specific care needs, with local services being required to support these needs in the most appropriate way. This was one of the key messages of Lord Darzi’s recent report “High Quality Care for All” and presents an exciting, yet challenging, framework in which care for long-term conditions must be developed.

Both these movements are good news for diabetes services and ultimately for all people with long-term conditions. But turning these changes into improved outcomes will not be easy. Shifts in service planning need to be made before the relationship between the person with diabetes and the service can change. It is this requirement that makes the Year of Care programme an ideal place to address these challenges, and is why the programme has captured the attention of so many people and groups.

When the pilot programme began, following a commitment in the 2004 White Paper *Choosing Health: Making healthy choices easier*, three healthcare communities in England were selected to pilot the Year of Care approach to both care and commissioning. Once implemented at a local level, the Year of Care rapidly grew from a discrete intervention into a complex programme of change, requiring services and organisational processes to be redesigned and starting a cultural shift amongst both healthcare professionals and people with diabetes.

The sheer scale of the programme is highlighted by the number of people who have become involved in it, both locally and nationally. Within the sites it has brought together Board members, consultants, GPs, diabetes specialist nurses, podiatrists, practice nurses, commissioners, Public Health, administrative and communications teams, Patient and Public Involvement leads and, of course, local people with diabetes.

Support for the national team has come from our partner organisations: Diabetes UK, The Health Foundation, the National Diabetes Support Team and also the Department of Health. The NHS Alliance and the NHS Confederation have been on
hand to offer advice, as has the National Association of Primary Care; we are indebted to the work of NHS Yorkshire and the Humber and have been fortunate enough to use Healthcare for London and the Care Planning Champions’ Network as sounding boards for ideas.

We hope that this publication will begin to pass on the knowledge gathered so far from the Year of Care. I hope you find it useful, whether it makes you stop and reflect about the care you provide, or inspires you to implement it locally. My only request is that you let us know of any work you undertake locally so that we can continue to share enthusiastic practice and celebrate local successes.

Rowan Hillson

Dr Rowan Hillson
National Clinical Director for Diabetes
Introduction

This guide to *Getting to Grips with the Year of Care* is designed to help commissioners, clinicians and networks interested in setting up Year of Care programmes locally.

The Year of Care programme sets out to learn how routine care can be redesigned and commissioned to provide a personalised approach, including support for self management, for people with long term conditions, starting with diabetes.

**It is a three year programme. The guide focuses on the preparatory phase only** (i.e. the first year). It concentrates on what needs to be put in place and particularly the key role commissioning has to play.

Each section provides guidance based on national literature and the experience of the three Year of Care pilots.

These pilots, in North of Tyne, Calderdale and Kirklees and Tower Hamlets, aim to:

- Establish care planning in routine practice.
- Identify sections of the local diabetes population by potential need for services and support for self care.
- Develop new and existing providers to support self care.
- Systematically link individuals’ needs and goals into population level commissioning.
- Explore the costs and benefits of providing these services and support.

The feasibility phase was successfully concluded in August 2008 and the evaluation of the implementation of the Year of Care was completed shortly after. The implementation phase will run through until 2010. An independent evaluation is assessing the impact of the approach.

Building on the experience of the pilots this guide:

- Outlines the thinking behind the Year of Care.
- Highlights the building blocks that need to be in place locally.
- Provides practical tips and advice on how to get started.

Key initial learning is that the services for people with long term conditions need to change fundamentally and this means that the Year of Care approach needs to be seen as a vast change project.

Since inception, the pilot sites have evolved with the Year of Care programme to become demonstration sites acting as laboratories for change. The good news is that as understanding of the scale of the task has got larger, motivation and commitment to local sustainability has become stronger. This has been largely based on the positive feedback from service users and clinicians, and the commitment to the patient at the heart of the endeavour.

Whilst we can offer resources, examples, a demonstration teaching programme and a core slide set that outlines exactly what the Year of Care is and its benefits, we cannot change local culture. That needs to be done by you. You will need considerable local understanding, enthusiasm and drive in order to put in place the changes required to deliver a Year of Care effectively. The Year of Care can be used as a catalyst for a programme of transformation. It therefore needs the commitment of an entire healthcare community.

We hope this guide is useful to organisations wishing to set up the Year of Care. The examples included are a small part of the material the pilots have produced, which is all available on the linked website. We have signposted these and other resources throughout the guide, and would always recommend that in the first instance you contact the Year of Care central team at yearofcare@diabetes.org.uk.
What can the Year of Care offer me?

The Year of Care has many stakeholders; there are potential benefits for everyone.

Benefits to Chief Executives and Boards:

- It provides the framework for personalised care in long term conditions outlined in the Next Stage Review.
- It provides the infrastructure for the 2010 commitment for everyone with one or more long term condition to be offered a care plan.
- Greater value for money, as services are provided which meet local needs and deliver improved health outcomes.
- A stimulus to the whole healthcare community to redesign services for long term conditions, ensuring the right care is provided in the right place at the right time by people with the right skills, with the right funds in the right place.
- Potential to reduce avoidable admissions and A&E costs.

Benefits to people with long term conditions:

- More involvement in planning their care.
- A better understanding of their condition.
- Respect for and recognition of their everyday work to self manage.
- Tailored support for those who want to take on a greater role.
- Consistency and continuity of care.
- A central role in service planning, and agreeing what local care should look like.
- Information and signposting to local support services.

Benefits to clinicians:

- More satisfying consultations.
- A lever with commissioners to agree more time with patients.
- Commissioning influenced by genuine clinical data.
- Services commissioned that people with diabetes will use.
- Better outcomes for people with long term conditions.
- A new and interesting skill set.
- A lever to improve clinical IT and drive quality improvement.

Benefits to commissioners:

- Complete fit with World Class Commissioning.
- Provides the information needed to commission services that people want and clinicians value.
- Increases effective self management and addresses local need.
- Has a positive impact on other local and national drivers, such as reducing acute admissions and improving the patient experience.
- Provides a long term, sustainable approach to reducing the burden of chronic diseases on local resources.
- A constructive environment to work with clinicians.
- A detailed understanding of pathways and costs as the basis of new local currencies.
**What is the Year of Care?**

Year of Care is about improving care for people with long term conditions in the NHS. It describes the ongoing care a person with a long term condition should expect to receive in a year, including support for self management, which can be costed and commissioned.

The Year of Care approach puts people with long term conditions firmly in the driving seat of their care and supports them to self manage. It does this firstly by making routine consultations between clinicians and people with long term conditions truly collaborative through care planning, and then ensures that the local services people need to support this are identified and made available through commissioning.

To achieve the best outcomes, both effective care planning and commissioning have to be in place and working together.

Each individual will have different priorities and goals, often leading to actions they can take themselves. They may also choose how these will be supported from a wide range of differing service and support options (micro-level commissioning). Many commissioners are already providing a range of self care support services to build on. The challenge is to link each individual’s needs and goals, choices and service use into the commissioning decisions that take place at population level (macro-level commissioning). If this can be achieved, services can be planned based on the genuine needs of individuals.

---

**Figure 1**

*Year of Care: Linking clinical care and commissioning*

- **Individual patient choices via the care planning process = micro-level commissioning**
- **MENU OF OPTIONS**
  - Information
  - Structured Education
  - Weight management
  - Screening for complications
  - Telephone review/support
  - Smoking cessation advice
  - Local authority exercise programme
  - Specific problem solving
  - EPP
  - Buddying / walking groups
- **‘An end in itself’**
- **‘A means to an end’**

*Macro-level commissioning by the commissioner (PCT/practice) on behalf of the whole diabetes population*
The Year of Care concept was originally linked to the idea of commissioning for periods of care in people with long term conditions using secondary care services, and was first referred to in this context by Professor Pieter Diegling. From the very start there have therefore been differing interpretations of what a Year of Care actually means. The three pilot sites interpreted the term in different ways, and our observation is that this will be the case in any healthcare community that begins to talk about it. Having different views within the team or in the healthcare community is an enormous barrier to delivery.

The Year of Care programme as it stands today, and the work of the pilot sites that is central to it, is about taking a new look at the relationship between the NHS and people with long term conditions; the services they need to support them in living with their condition; adapting to and owning the changes it brings; and finally to becoming more effective in their self management of it. On the way the programme will collect a great deal of information about the services people use in a year from all parts of the healthcare community, and what this costs.

The information and resources discussed in this Guide relate only to the definition of Year of Care given on the previous page.

One of the most important first steps for any community wishing to start thinking about Year of Care is to understand and agree what they mean by it.

Common areas of confusion

Many commissioners in England are currently looking intensively at how diabetes services are organised in their communities and coming up with new models of care, new roles for staff and new pathways for patients. It is natural for them to assume that this is what Year of Care refers to.

While work focussed on coordinating services is very important, the Year of Care programme and approach is focussed on rethinking the connection between the person with the long term condition, healthcare professional and the service.

It follows that the Year of Care is not about describing new models or pathways of care. However, all three of the Year of Care pilot sites that started out in 2007 concentrating solely on core commissioning tasks for the Year of Care itself, have found that they cannot take full advantage of the benefits - nor sustain it in their communities - without addressing the underlying model of care as well. These aspects and how to get to grips with them are described in the section on commissioning.

Healthcare communities which plan to introduce the Year of Care will find it helpful to look at their model of care and wider commissioning strategies right from the start.

Finally, the Year of Care is not a single intervention that will either work or not work, and it is not being rolled out nationally. The Year of Care is a response to the huge challenges facing care of people with long term conditions, and provides a learning environment and community of practice which we are inviting people to join.
The involvement of people with long-term conditions in planning their own care and choosing how to manage their own condition is a critical step towards improving patient related outcomes and the focus of national policies to improve quality, such as the National Service Framework for diabetes (1).

However, the evidence is that we still have a long way to go to achieve that aim. In 2007 the Picker Institute reported that whilst NHS care has improved markedly in some respects (e.g. waiting times), and most patients are highly appreciative of the care they receive, the service as a whole is still far from patient centred (2). The most significant problem is that patients are not encouraged to share their views and feel they do not receive enough help to effectively self care.

Why a Year of Care?

In 2007, the Healthcare Commission published their review of almost 70,000 people with diabetes in England (3). They found that although 95% of people had diabetes checks at least once a year, less than half discussed their goals in managing their condition (Figure 2) , although having clearly articulated goals is the most important link with changing health behaviours.

Diabetes is also one of the most complex long term conditions. It already affects nearly 5% of the population, uses nearly 10% of NHS resources and contributes to 10% of deaths. Prevalence is increasing fast. Diabetes can affect almost every area of the body and impinges on nearly every area of NHS activity.

So the need is great and the challenge considerable. If the Year of Care can provide personalised and tailored services for people with diabetes, the lessons learned and the new skills can be transferred more easily to other long term conditions.

Why a Year of Care for People with Diabetes?

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Figure 2

Diabetes care: The percentage of adults who report that they…

- had at least one check up in the last 12 months
- and discussed ideas about the best way to manage their diabetes
- agreed a plan to manage their condition over the next 12 months
- discussed their goals in caring for their diabetes

“We have to keep up with the expectations of the public. This will mean allowing people to exercise choice and be partners in decisions about their own care, shaping and directing it with high quality information and support”.

“The fundamental solution to the rise of lifestyle diseases is to change our lifestyles. While the NHS can support and encourage change, ultimately, these are decisions that can only be made by us as individuals. Those with two or more long term conditions are more likely to be obese, eat less healthily and smoke than those with one or none (5). People need to know the risks and have the opportunity to take control of their own healthcare”.

“International best practice suggests that control by a patient is best achieved through the agreement of a personal care plan (6). Care planning creates packages of care that are personal to the patient. It involves working with professionals who really understand their needs, to agree goals, the services chosen, and how and where to access them. Personal care plans are agreed by the individual and a lead professional”.

“Over the next two years, every one of the 15 million people with one or more long term conditions should be offered a personalised care plan, developed, agreed and regularly reviewed with a named lead professional from among the team of staff who help manage their care”.

“Primary care trusts and local authorities have the responsibility to ensure that all this is achieved, as well as offering a choice of treatment setting and provider”.

Lord Darzi’s review and the clinician led SHA reports on which they were based were themselves drawing on a wealth of previous thinking and policy. Care planning was a central component of the Diabetes National Service Framework (1) and resulted in the report of a Working Group in which people with diabetes and clinicians defined what needed to be done. This was developed further for all people with long term conditions and tools for commissioners provided (7).
The Year of Care is a change project which links across the health and social care communities. It is not a quick fix for personalised care, but requires sustained action and so ensuring that everything is in place from the outset is absolutely critical to its success. From phase one of the project we have learnt that there were three important initial issues.

1. Commitment from the top of each pilot organisation with sustained involvement from a committed and enthusiastic Senior Responsible Officer, a formal steering group and employment of a full time project manager was critical to the huge amount of work and positive outcomes achieved in the first nine months. Pilots found that one month from successful application to start up was not enough to secure this effectively but longer than three months would have reduced the focus.

2. The project needs to be an integral part of the organisation’s commissioning agenda, and synergies and tensions with other parts need to be managed.

3. Developing a clear local understanding of what the Year of care means should be an early priority.

Not all the sections below will be relevant to everyone. Some organisations or services will find that they are further ahead than they thought.

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**Figure 3**

**Setting up the Year of Care will be easier if there is:**

- High level clinical and managerial buy-in.
- Board level sponsorship.
- Local engagement and enthusiasm.
- Developed partnerships between voluntary, health, social care and private sectors.
- Strong local leadership in diabetes care.
- Commitment to people development and involvement.
- Robust management processes.
- Results oriented culture, with outcome measures in place.
- Active involvement of users.
### Key messages from the pilots

1. Recruit a Project Manager.
2. Agree clear governance structures – who is accountable for what, who is leading each stream, who has overall responsibility.
3. Develop a project team after consideration of both functions and key individuals; take time to meet regularly, perhaps via away days and workshops.
4. Identify local ‘champions’ – ensure these are both clinical and managerial. Senior clinicians will need to devote considerable time which needs backfilling and this can take a few weeks to organise.
5. Agree clear reporting mechanisms.
6. Agree clear channels of communication.
7. Agree a communications strategy – how to promote the Year of Care internally and externally, and how to feed back results to people who have contributed. Linking with the work of the diabetes network will be important, to harness skills and avoid duplication.
9. Agree local support mechanisms.
10. Secure local commitment.
11. Identify resources needed – time, staff, expertise.
12. Set up programmes of work to deliver the required infrastructure for the care planning ‘house’ and commissioning ‘windmill’ models (see below).
13. Agree a workable timeline, with appropriate milestones.
Stakeholders
The following is a compilation of stakeholders included in pilots. This should not be considered as a checklist as individual PCTs will identify their most relevant and important local stakeholders. Since a majority of the new work will take place in primary care where 85% of clinical contact in diabetes takes place, having senior and enthusiastic involvement from primary care is essential and this will need to be resourced. People with cross organisational roles were found to be particularly useful.

Figure 5

<table>
<thead>
<tr>
<th>Project Team Stakeholders</th>
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<tbody>
<tr>
<td>• Senior Responsible Officer</td>
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<tr>
<td>• Project manager</td>
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<tr>
<td>• Medical Director</td>
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<tr>
<td>• Clinical diabetes lead</td>
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<tr>
<td>• Commissioning lead</td>
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<tr>
<td>• Managerial diabetes lead</td>
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<tr>
<td>• Diabetes physician</td>
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<tr>
<td>• GP lead</td>
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<tr>
<td>• Clinical psychologist</td>
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<tr>
<td>• Public Health lead</td>
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<tr>
<td>• People with diabetes</td>
</tr>
<tr>
<td>• Dietician</td>
</tr>
<tr>
<td>• Podiatrist</td>
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<tr>
<td>• Social care commissioner</td>
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Figure 6

<table>
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<tr>
<th>Other Stakeholders</th>
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<tbody>
<tr>
<td>• GP</td>
</tr>
<tr>
<td>• Practice nurses</td>
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<tr>
<td>• Practice managers</td>
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<tr>
<td>• Practice administrators</td>
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<tr>
<td>• Receptionists</td>
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<tr>
<td>• Healthcare Assistants</td>
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<tr>
<td>• Practice based commissioning chairs</td>
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<tr>
<td>• Primary care diabetes leads</td>
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<tr>
<td>• Community nurses</td>
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<tr>
<td>• District nurses</td>
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<tr>
<td>• Secondary care consultants</td>
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<td>• Podiatrists</td>
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<td>• Dieticians</td>
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<tr>
<td>• Ophthalmologists</td>
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<tr>
<td>• Pharmacists</td>
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<tr>
<td>• Director of nursing</td>
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<tr>
<td>• Community matrons</td>
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<tr>
<td>• Long term conditions board members</td>
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<tr>
<td>• Continuing Professional Development leads</td>
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<tr>
<td>• Public Health colleagues</td>
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<tr>
<td>• Information analysts</td>
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<tr>
<td>• Patient and Public Involvement Forums</td>
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<tr>
<td>• People with diabetes and service user groups</td>
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<tr>
<td>• Patient advice and liaison service coordinators</td>
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<tr>
<td>• The public</td>
</tr>
<tr>
<td>• Diabetes UK patients/supporters</td>
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<tr>
<td>• Voluntary sector services e.g. exercise groups</td>
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<tr>
<td>• Healthy Living Centres</td>
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<tr>
<td>• Public libraries</td>
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<tr>
<td>• Adult social care</td>
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<tr>
<td>• Education and training providers</td>
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Pilots’ Experience
Each pilot site developed a primary care engagement strategy. This varied according to local culture and previous work. Care planning had already been piloted in North of Tyne and the benefits had spread by word of mouth between practices. In Calderdale and Kirklees, the project manager had worked with practices previously and had already developed excellent working relationships. In Tower Hamlets, a senior and highly regarded GP took on the role of champion and tried out many of the new features in her own practice first.

Other effective approaches included:
- Informal ‘meet and greets’ involving senior Trust staff and primary care.
- The sharing of all possible benefits with staff.
- Use of bite size slots at local meetings, such as GP and nurse meetings.
- Short articles included in local newsletters.
- Involving practice managers and administrative staff from the outset.
- Valuing the local knowledge and expertise of local practices to inform commissioning.

North of Tyne used a two staged approach to engage primary care.

1. The commissioning lead visited practice based commissioning groups to share project information and to get provisional sign up.
2. An information pack was then produced and distributed to practices with an invitation for the diabetes lead GP and practice nurse to attend Year of Care training days.

This approach proved successful, with 29 practices out of 30 from across North Tyneside signing up.

Key guidance to support commissioners and their networks
- The National Diabetes Support Team has produced guidance for project leads and project team members to help develop project plans and to achieve project aims, Diabetes Service Planning: A Project Management Guide (8). The guide includes helpful tips and templates which can be adapted to meet local needs.
- A presentation outlining the Year of Care concept and its benefits can be obtained by emailing: yearofcare@diabetes.org.uk

Key guidance to support stakeholder engagement
Examples of the information packs produced by the pilots are available on the Year of Care website:
One of the most useful things that the pilot sites developed was a clear vision of the whole of the Year of Care project, and then a series of images and models which broke this down into the many practical tasks.

The key images are the care planning house and the commissioning windmill and this guide is designed round their constituent parts. Different stakeholders will be involved in different aspects of the project but the essential thing is that everyone can see the ‘bigger picture’ and where their contribution fits in, because the parts are all interconnected.

This ‘bigger picture’ is the care planning house model, which is built on the foundations of robust commissioning processes. It demonstrates that the crucial contributions of an engaged, informed patient and a healthcare professional committed to partnership working together also need an appropriate environment with sound organisational processes that facilitate their interaction.

The ‘house’ metaphor emphasises the importance and inter-dependence of each element - if one is weak or missing the structure (i.e. the diabetes service) is not fit for purpose.

**Care Planning and Commissioning in Practice**

![Organisational processes diagram](image)

- **Engaged, informed patient**
- **The care planning consultation**
- **Commissioning - The foundation**
- **HCP committed to partnership working**

HCP - Healthcare Professional
Care planning can be seen as an end in itself. The value of engaging people with diabetes, or with any long term condition, in their healthcare and self care components is supported by a wealth of evidence. It is at the heart of partnership working. The National Diabetes Support Team’s publication, ‘Partners in Care: A guide to implementing a care planning approach to diabetes care’ (9) describes it as: “Looking with rather than at someone with diabetes”

In the context of Year of Care, care planning is also a means to an end. The care planning consultation will identify what each individual needs to support them to self care effectively. Across a health community these elements can then be collated and used by commissioners to ensure that the specific support and services required are being provided and available to use. Care planning will also be the essential portal to access many personal care opportunities such as individual budgets.

It’s a verb!
Perhaps the most important message for everyone involved in care planning is that while having a written care plan is important for many people, it is the process of collaborative care and joint working rather than the end product which is helpful. It is the action, ‘care planning’ which is important not the product, the ‘care plan’. It is the verb rather than the noun!
The National Diabetes Care Planning Working Group has developed a model for effective care planning (10) (Figure 8), based on the vision set out in the Diabetes NSF and the Modernisation Agency’s report on care planning (7).

Figure 8

The care planning consultation

The National Diabetes Care Planning Working Group has developed a model for effective care planning (10) (Figure 8), based on the vision set out in the Diabetes NSF and the Modernisation Agency’s report on care planning (7).
This sequence of gathering, sharing and discussing information, deciding the issues, and developing a plan is not too distant from the traditional clinical consultation. However, in care planning the patient views and a much wider range of issues across all aspects of a person's life with their condition is taken into account, and their personal goals are central to the action plans developed. In diabetes it can replace the traditional annual review which has sometimes become a ‘tick box’ activity, with a ‘conversation’ that is more satisfactory and effective for everyone. The components are discussed in detail in *Partners in Care* (9).

### Figure 9

**The elements of the care planning consultation**

1. **Gather and share stories:** The process begins with the equal sharing of both the professional’s and patient’s stories or perspectives. For the person with diabetes, this should start before the consultation with active encouragement to reflect on their life with diabetes, and questions or concerns they may have and would wish to raise in the consultation. In addition, this could include sharing of biomedical results prior to the consultation.

2. **Systematic review of domains:** Just as a traditional clinical ‘history’ has a structured approach to eliciting symptoms and signs, a care planning approach can review the ‘domains’ that are important to managing a long term condition like diabetes. The clinical domain is obvious but the domains of emotional health, social and behavioural issues and tapping into knowledge and health beliefs are equally important in supporting individuals to self care, but may often be overlooked.

3. **Exploring and discussing information:** This is central to a true partnership and a necessary prerequisite to deciding on the next stage, the individual priorities and more importantly their goals.

4. **Goal Setting:** This critical component, in which the earlier stages feed into a focus and identification of an explicitly stated individually determined goal, is the step which the literature tells us is most closely associated with subsequent effective self management.

5. **Action planning:** At this stage of the consultation, the person with diabetes and the healthcare professional should agree a set of action points, decide who will be responsible for achieving each of the actions and agree when the actions will be reviewed. For the person with diabetes, the actions are likely to relate to aspects of self care (9). For the healthcare professional, they might include referrals (especially to the self support services listed in the commissioner’s menu of options), investigations etc. Actions are more likely to be undertaken by either party if they are detailed, specific and set out within a given timescale.
The components of Care Planning

Figure 10

The pilot sites have found that there were a variety of ways to support the key components of the ‘house’, which can be broken down into the various ‘building blocks’ shown below. The text around the house shows examples of the key issues that need to be considered under each ‘block’.

The next sections look at what is required for strong ‘walls’ and a secure ‘roof’.
Engaged, informed patient

To enable an individual to self-manage effectively and participate fully in any decision making processes, they will need a good understanding of their condition and how they can access any additional information they need. They also need to be familiar and comfortable with their potential role in partnership with the healthcare professional in the consultation and decision-making.

All sites see the benefit of commissioning structured patient education courses as a core part of routine care for people with diabetes to ensure that they are able to take part in care planning with more confidence and information. It is important to recognise that there are other mechanisms and sources of information, including ensuring that people are offered access to their records, or receive copies of clinical correspondence written about them (11).

A first step

The sharing of results and other information prior to the consultation has been piloted and successfully implemented in a number of settings in both primary and specialist care in North Tyneside and Northumberland, and has been highly valued by both patients and health care professionals (9). The common components of the process are outlined in Figure 11. This process has been successfully reproduced by practices in each of the pilot sites with positive results and feedback.

This aspect can be started quite quickly by clinical teams and provides positive motivation to engage with other aspects.

Figure 11

- Information gathering
- Information sharing
- Care planning consultation
- Agreed and shared care plan
Other aspects of communication with patients

People with diabetes need information about the care planning process itself, what to expect in this new way of working and how they can be involved.

It is essential to find ways of engaging and informing people with diabetes who have difficulty in reading and writing English, or whose first language is not English. For example:

- Narrative or picture based education programmes may need to be offered.
- Using telephone contact or recorded letters, rather than written, for reminders (both to provide information and for the care plan).
- Use well trained health workers from similar ethnic backgrounds to act as advocates where necessary.
- Multilingual interpreters have been shown to be very effective where employed and used appropriately.

Where there are populations with low health literacy extra resources may be needed initially to ensure that people understand the material and how to use it. These issues were explored by Tower Hamlets and examples are available by contacting yearofcare@diabetes.org.uk.
Healthcare professionals committed to partnership working

Engaging Local Staff

True partnership working will require a different approach to the consultation for many healthcare professionals. However, the following comments are frequently heard:

• Providers are not committed to this way of working.
• People with diabetes do not want this approach.
• You cannot measure this approach effectively.
• Working like this takes up too much time.
• People disagree with your care plan.
• Doing care planning means we will lose Quality and Outcomes Framework (QOF) points.

A suggested approach to all these issues and more is available in ‘Partners in Care’ (9).

Contrary to being sceptical, a further issue is that many clinicians will say, and believe, that they do this already. For those in primary care, a patient centred approach to consultation is at the heart of modern training. The evidence presented earlier is that for patients who want to be increasingly in control of their long term condition, this is not enough. It does however provide an excellent basis on which to move on and incorporate the new and additional skills many practitioners will need, to support self care and be comfortable with the patient being in the driving seat of their care.

The demonstration training programme (see Appendix I) is set up to enable these discussions to come to the fore, to provide situations in which to debate them and enable practitioners to reflect on their own practice. This programme is modelled on and mimics the new approach to care planning in which practitioners can set their own goals and action plans for improving their personal practice. Tools are available to help them assess their own practice. The use of consultation quality measures has been piloted, as well as team based assessments of the effectiveness of the clinic organisation in supporting self care.

Pilot sites have found that in addition to providing training there is no substitute for shoe leather. Seconding senior staff and using people who are already known and trusted in primary care have been essential to engaging and supporting practices in the tasks they need to do. Once started the positive feedback from patients and the increased satisfaction with more productive consultations help to generate further momentum.

Reviewing and Meeting Staff Training Needs

The pilot sites all developed or commissioned training programmes for clinical teams. They pooled their experience and developed criteria for success. North of Tyne, which had the largest group of practices of the three sites, also had expert educators as part of their core project team. They were able to test, modify and repeat their training, and working with the other sites this has become the core of a demonstration programme (see Appendix). This complies with the national criteria that all education programmes should meet.

Key learning points for Year of Care sites when thinking about training include:

• Training should be endorsed at the organisational level by senior leaders.
• Each organisation should assess staff learning needs and use this as the basis for the training programme.
• Training should be multidisciplinary, including practice managerial and administrative staff, and health and social care professionals working in the local community.
• Trained trainers and senior clinicians should deliver the programme.
• Methods for maximising primary and secondary care professionals’ attendance should be used. Practices should be supported to release staff. This includes providing cover for staff attending training, and delivering training at times that do not impinge on clinical commitments, e.g. lunch times and evenings.
• Ensure new staff can access training.
• Do not underestimate the time and resources required for planning, coordinating and delivering the programme.

Content of training should include:
• What is Year of Care and key components.
• Philosophy of care planning and self care.
• Care planning consultation skills.
• Organisation and system aspects of care planning.
• Awareness of non-traditional services and support (menu of options).
• Use of local templates.
• Clear rationale for extra recording of patient information.

Embedding new skills
To embed and sustain the learning, training needs to be followed up with further support for practices.

Mechanisms to deliver ongoing support include:
• Practice visits.
• Small network groups.
• Locality level meetings.
• Learning sets.
• Care planning away days.
• Refresher sessions.

Key Guidance to Support Commissioners and Clinicians
The Year of Care and care planning training package is available in Appendix I
Organisational Processes
As the ‘house’ metaphor implies (p15), the effectiveness of the care planning consultation does not just depend on the quality of the interactions between the healthcare professional and the person with diabetes but also on the organisational processes that are involved.

On a simple level the usual registration, recall and review system will need to be adapted to allow the collection and sharing of the biomedical results prior to consultations. This will impact upon the wider practice team and their engagement and support is required. For example:

- Practice administrators will need to send out
  - Letters inviting people to the first appointment and explaining the care planning process.
  - Letters containing and explaining biomedical results and inviting people for the care planning appointments.
- A member of the clinical team, e.g. a Healthcare Assistant, needs to perform the biomedical tests prior to the annual review care planning visit.

In addition, mechanisms need to be in place to ensure the practice staff are fully aware of the menu of options or services that are available and how to access these.

Pilots’ Experience
As well as the resources required to implement these processes there are very practical implications. For instance, Tower Hamlets has produced folders for patients to store test results and care plans. Each folder contains a simple colour coded chart that helps patients to interpret their biomedical results. Service users highly valued the coloured charts that helped them understand their results, but colour printing was not available in all practices so appropriate printers had to be provided.

Calderdale and Kirklees have produced a results sharing sheet, ‘Getting the most from your appointment’, which encourages patients to set out the agenda they would like to discuss during their consultation, and a care plan pro forma.

The pilots have worked in partnership with service users and healthcare professionals to review all resources, including leaflets, posters and patient letters to ensure they are in line with Year of Care principles.

Key guidance to support commissioners and clinicians
- Examples of template letters, folders, information sheets, posters and care plans produced by the pilots are available on the Year of Care website: http://www.diabetes.nhs.uk/work-areas/year-of-care
IT Templates – Linking Clinical Care and Commissioning

While IT cannot make people change their behaviour, it can support change and make it easier for people to do the right thing. Many GP practices are now ‘paperless’ and pilot sites quickly found that once they knew what they wanted to do, their current systems were not geared to support this. New templates are needed not just to send out the right letters and results in the new style, but to record the new information from patients and the important personal goals and action plans at the heart of support for self care.

Practices also need to have a system for capturing information from the care planning consultation, particularly goals and actions, to feed into macro-commissioning.

As an example, when considering the support or services provided for people who are overweight PCTs will currently know:

- The number of people who have a Body Mass Index (BMI) greater than 30.
- The services available to support people to lose weight.
- The capacity of each of these services.
- The number of people who use each service.
- The number of people who lose weight.

In addition, through the care planning process, Year of Care will help to:

- Identify the total number of people who want to lose weight.
- Identify the number of people who wish to access existing support services. This will also identify over and under capacity of existing services.
- Identify the other actions and interventions, including self help, that people would like to use but are not available in the current menu of options.
- Provide feedback on the effectiveness of the different interventions and support services.

The pilots have collaborated with an NHS Yorkshire and the Humber IT project, which has developed standardised templates that capture all these features. This is being tested initially in Yorkshire and will be used by the pilot sites in the near future. The template has been developed and is being tested using TPP SystmOne (see Appendix II). The templates and specification have been made available for incorporation or adaptation into other systems but this may not be a simple process.

Sites thinking about exploring the Year of Care need to give thought to this early on as the lead in to get the changes needed may become a rate limiting factor.

Pilots’ Experience

The pilots have shown that it is possible to implement simple templates and systems to allow the results sharing letters to be produced relatively easily. They have also attempted to make a start using manual systems for capturing and analysing the outputs of the care planning process for commissioning purposes.

Key Guidance to Support Commissioners and Clinicians

- Further information on the NHS Yorkshire and the Humber IT project can be obtained from:
  James Thomas
  Year of Care Programme Manager
  Healthcare Policy Team
  Diabetes UK
  Tel: 020 7424 1113
  Email: James.Thomas@diabetes.org.uk
The pilot sites initially identified four specific commissioning tasks to support the Year of Care Programme:

1. Commissioning care planning.
2. Developing the menu of local options.
3. Linking micro-level to macro-level commissioning.
4. Service user involvement.

As work progressed they found that these specific elements linked to the wider work they were undertaking across their local organisations, within the framework of World Class Commissioning.

Commissioning for the Year of Care thus became a highly complex endeavour with many component parts. As with care planning, pilots found it was more manageable if broken down into a set of linked tasks and work areas.

These can be described as the sails of a windmill, an image of sound delivery firmly based on the generic principles of World Class Commissioning. Each sail represents an area of relevant commissioning activity. The central part of each sail identifies one of the specific work areas of the Year of Care without which it will not be possible to deliver the benefits. The outer part of each sail is the linked activity the pilot sites found they were drawn into, both to enhance the benefits and to ensure sustainability and local spread.

Understanding these links and establishing Year of Care within a wider strategic commissioning strategy would be helpful for those embarking on this journey.

Figure 12

<table>
<thead>
<tr>
<th>World Class Commissioning Competencies</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Locally lead the NHS</td>
</tr>
<tr>
<td>2. Work with community partners</td>
</tr>
<tr>
<td>3. Engage with public and patients</td>
</tr>
<tr>
<td>4. Collaborate with clinicians</td>
</tr>
<tr>
<td>5. Manage knowledge and assess needs</td>
</tr>
<tr>
<td>6. Prioritise investment</td>
</tr>
<tr>
<td>7. Stimulate the market</td>
</tr>
<tr>
<td>8. Promote improvement</td>
</tr>
<tr>
<td>9. Secure procurement skills</td>
</tr>
<tr>
<td>10. Manage the local health system</td>
</tr>
<tr>
<td>11. Make sound financial investment</td>
</tr>
</tbody>
</table>
The Year of Care commissioning 'sails'

1. The care planning sail

Year of Care specific task
Care planning itself needs to be commissioned together with other elements of the ‘house’. Some of the issues that will need to be considered are:

- How can this be embedded contractually? Does this need to be rewarded, perhaps via a Local Enhanced Service, or will it be included in General Medical Services?
- Potential increases in cost, and increases in time per consultation, at least to begin with. How will the expected savings from reductions in follow on appointments and reduced use of hospital resources be recorded?
- Ensuring healthcare professionals have the appropriate skills and that training is available and taken up. How will new skills be supported and maintained? How to commission high quality training that meets the objectives of their programme. How robust are existing peer review structures, and what can be done to ensure that this is undertaken routinely?
- How will delivery of a care planning approach be monitored?
- How can a consistent approach to consultations be guaranteed to patients?
- Are local IT systems and Electronic Health Records capable of supporting care planning, and trapping appropriate commissioning data?

Pilots’ Experience
North Tyneside PCT is currently revising its local enhanced service to recognise the changes that are required to implement effective care planning.

Kirklees PCT has developed a service level agreement to encourage practices to produce a plan for implementing care planning with their patients.

Tower Hamlets has had a local enhanced service specification for practice based patient centred diabetes services for some years. The Year of Care programme has identified a need for increasing the tariff paid to pilot practices to address the increase in resources required. Additional resource may be needed initially to support people with poor health literacy, who are keen to participate fully in the care planning process.

Resources
The central team has made available:

- A demonstration specification to commission care planning services.
- A demonstration training curriculum. A quality assurance tool to enable commissioners to assess training that they may be procuring will be available soon.
- IT templates to record the care planning consultation, patient use and service need and use.
- Tools for organisational audit and consultation evaluation (CQI-DM).
The Wider Commissioning Tasks

Year of Care raises many issues for the whole local diabetes service. Among these are:

• How does care planning fit within the local model of care? Is there a systematic approach to where people with diabetes are seen?

• What is the role of the local specialist service?

• Is structured patient education incorporated into local pathways as routine, and does it specifically link to care planning itself?

• Are the resources in the right place to support the Year of Care Programme?

• How will funding be moved in line with any movements in care provision?

• What does this mean for local workforce development plans?

North of Tyne recognised the importance of this at an early stage and the potential for the Year of Care project to be a local lever for change. They have carried out an in depth piece of work across their whole commissioning area using programme budgeting and marginal analysis. This identified considerable variations in expenditure and outcomes and has enabled them to take a structured approach to redesigning services with care planning and a Year of Care approach across a much wider level.

All the pilots are now undertaking some form of structured approach to commissioning a redesigned diabetes service.

Designing the Model of Care

There is considerable advice and experience available in developing or redesigning a whole community wide diabetes service. The ‘model of care’ defines all the components of the service, including the workforce, which should be commissioned to ensure that local people have a service that fits their specific needs. The ‘Diabetes Commissioning Toolkit’ (12) outlines the components of a diabetes health needs assessment, and of a generic specification based on national quality standards including National Service Frameworks and National Institute for Health and Clinical Excellence (NICE) guidance. ‘Developing a Diabetes Service: Using the diabetes commissioning toolkit’ (13) provides practical suggestions and guidance.

It is important to maximise all existing local knowledge and expertise. As part of World Class Commissioning, organisations will already be looking to work with local clinical teams and groups. Where diabetes networks exist these provide an excellent place from which to base this work. Many networks are equipped to not only offer advice to commissioners, but also to share previous projects and plans which would fit well with specific elements of the Year of Care commissioning agenda. There is practical advice on how to include users in local networks.
Key Guidance to Support Commissioners and their Networks

Needs Assessments and Service Models
- Diabetes Commissioning Toolkit (12) to help NHS commissioners to adopt a strategic approach to developing diabetes services.
- Developing a Diabetes Service: Using the diabetes commissioning toolkit (13)

Patient Education

National Reports

Online Resources
- The National Diabetes Support Team and the Yorkshire and Humber Public Health Observatory have been working together to develop an online Diabetes Data Directory which:
  - Provides a brief description of the various tools/datasets.
  - Links directly to relevant websites.
  - Signposts to the various tools/datasets that will provide answers to key headline questions.

http://www.yhpho.org.uk/diabetesdatadirectory/introddd.asp
- A wealth of resources exists to support the process of mapping. Examples include: improvement leaders guides and process mapping, analysis and redesign guides. These can be found on the NHS Institute for Innovation and Improvement's website:

- Examples of booklets produced by the pilots containing a menu of options to support patients with self care are available on the Year of Care website:

2. The needs assessment sail

Assessing individual needs through the care planning process needs to be linked to wider approaches to local needs assessment including those within the framework of local Joint Strategic Needs Assessment.

Specific Year of Care Task
The pilots worked actively to identify the information they needed from routine clinical records (micro commissioning) to enable the link to be made with macro commissioning. This resulted in the development of Year of Care IT templates by NHS Yorkshire and the Humber.

Pilots’ Experience
The pilots recognised that other methods of needs assessment would be needed during the first year to inform the scope and scale of provider development.

North of Tyne piloted care planning and paper based recording in a number of practices.

Both Calderdale and Kirklees and Tower Hamlets used social marketing approaches leading to stratification of the population into categories with specific support needs.

Calderdale and Kirklees did a systematic review of diabetes records in one practice. This required significant human resource, and was to some extent hampered by the traditional biomedical content of the records. All sites held focus groups, learning from each other on how to run these and providing learning for others. The outcomes of Calderdale and Kirklees’ focus groups are shown in figure 13.

Wider Issues
There was considerable overlap with other needs of the populations, for example concerning obesity and Cardiovascular Disease (CVD) services. In addition, many patients had more than one condition (see the provider development sail, p32). Issues that pilots need to address included:

3. How to ensure your sample is representative of the local population.
4. How the data from the Year of Care fits with other short-, medium- and long-term local intelligence.
5. How IT templates can be developed to capture data routinely, and how this can be interpreted.
6. How patient confidentiality can be managed within the Data Protection Act and local operating guidance.
<table>
<thead>
<tr>
<th>Theme</th>
<th>Service User Preferences</th>
</tr>
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<tbody>
<tr>
<td>Emotional Support</td>
<td>• Someone to talk to/a buddying system/peer support</td>
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<tr>
<td></td>
<td>• Support for families and carers</td>
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<td></td>
<td>• 24 hour telephone helpline with trained staff</td>
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<td></td>
<td>• Less busy staff with time to listen</td>
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<tr>
<td></td>
<td>• More recognition, help and support for depression</td>
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<td></td>
<td>• Massage and meditation</td>
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<tr>
<td>Access to Services</td>
<td>• Closer to home</td>
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<td></td>
<td>• One stop shop, e.g. pharmacists offering phlebotomy</td>
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<tr>
<td></td>
<td>• Do not want to attend hospital for blood tests</td>
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<td></td>
<td>• Weekend/evening clinics with eye specialists</td>
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<td></td>
<td>• Easier/more access to foot services</td>
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<tr>
<td></td>
<td>• Longer opening hours with drop in clinics</td>
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<tr>
<td></td>
<td>• Professionals to keep to appointment times</td>
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<tr>
<td></td>
<td>• Professionals to follow up on what they say they will do, e.g. telephone contact</td>
</tr>
<tr>
<td>Information</td>
<td>• Clearer communication of results with an explanation of what they mean</td>
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<tr>
<td></td>
<td>• Innovative use of technology, e.g. test results via email</td>
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<td></td>
<td>• Information available in different languages</td>
</tr>
<tr>
<td></td>
<td>• Information for people with disabilities, e.g. deaf/blind</td>
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<td></td>
<td>• High quality information throughout care, not just at the beginning</td>
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<tr>
<td></td>
<td>• More advice on food and weight management</td>
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<tr>
<td></td>
<td>• A plan of what to expect in the next 12 months</td>
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<td></td>
<td>• More group education evenings</td>
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<tr>
<td>Lifestyle/social aspects</td>
<td>• Dosette boxes provided free or charge</td>
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<tr>
<td></td>
<td>• A greater range of activities to support a healthy lifestyle, e.g. t’ai chi, yoga</td>
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<tr>
<td></td>
<td>• Subsidised private gym membership</td>
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<tr>
<td></td>
<td>• Subsidised weight loss classes</td>
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<tr>
<td></td>
<td>• Training for staff in leisure centres</td>
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<tr>
<td></td>
<td>• More emphasis/guidance from doctors and nurses on non-medical lifestyle activities</td>
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<tr>
<td></td>
<td>• Single sex groups for some people</td>
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<tr>
<td></td>
<td>• More local support groups to encourage lifestyle change</td>
</tr>
</tbody>
</table>
3 The provider development sail

The way in which service providers can be supported to develop an appropriate menu of services to support self-management needs to be considered within the wider long term conditions agenda.

Some of the issues that need to be considered are:

- How social marketing techniques can be best employed to stimulate innovative providers.
- How the relationships with service providers can be effectively managed through any subsequent change processes, including decommissioning.
- How your local market can be formally developed.
- How to ensure that healthcare professionals are aware of current and future services.
- How to engage with and feed into existing long term conditions’ alliances.
- How to develop currencies for individual services, packages of care and entire long term conditions.
- How to work in partnership with local authority and health and social care colleagues, to develop tailored services.

Further learning will emerge over the next phase as pilots begin to implement their programme.

This needs to extend beyond the diabetes services because of the considerable overlap with long term conditions. For example, many of the services for people with diabetes would also be useful for those with cardiovascular disease, obesity or arthritis. These services are likely to be spread across the public, voluntary and private sectors.

As a first step, the commissioner will need to review the current services that support self care. This needs to extend beyond the diabetes services because of the considerable overlap with long term conditions. These are likely to be spread across the public, voluntary and private sectors. These might include, for example, prescription for health at a local gym, alternative medicine care, community group programmes, counselling services, citizens’ advice services, local patient and carer support groups, expert patient programmes and the PCT’s smoking cessation service.

Pilots found that not all these services were known to local practices. Some could be extended to fill gaps in provision identified in the local needs assessment and added to the menu of diabetes support services.

Pilots also set about systematically considering new providers and market development. Two pilots used social marketing techniques to segment the population into groups with specific needs so that these could be commissioned. This produced some debate within the project on the potential conflict of this approach with the care planning philosophy of building on individually identified goals and the potential for the service to ‘label’ individuals rather than groups. However, this approach provided new and imaginative ideas for support at population level and the potential to commission new services for the first year of the project delivery.

Using the Feedback from the Pilots

In Calderdale and Kirklees potential providers from across the health service, social services, local authority and the voluntary sector were invited to a workshop to map the current and planned Year of Care menu of options. Two groups reviewed existing levels of provision and identified coverage, gaps and overlaps between what was currently being provided or planned and what is actually needed. One month later, the group reviewed proposals to fill identified service gaps from potential providers.
The person with diabetes is the focus of the Year of Care programme, and must be a key player in its strategy, design and delivery. Throughout the set up phase people with diabetes, and their feedback, led to substantial learning and important changes in all areas of the project. People with diabetes were part of the steering groups in each pilot site, the national programme, pilot selection and evaluation teams and took part in many of the learning sets. The Year of Care benefited from being linked to wider Patient and Public Involvement and Health Equality programmes and this needs to occur right from the start. This was most successful where the different commissioning strands worked closely together, and in Tower Hamlets the Patient and Public Involvement lead was part of the Year of Care project management group.

Consulting Service Users Locally
Most activities have worked two ways, gauging attitudes to and opinions of the Year of Care concept and at the same time identifying concerns with existing service provision and potential solutions.

A variety of ways of involving users were employed such as focus groups, surveys and user reference groups. An approach that is useful for one purpose in one community may not work for another and pilots valued sharing experience of what was successful.

Traditional methods that might be used include:
- Advertising in local media.
- Personalised letters sent from practices.
- Links via established voluntary sector organisations.
- Use of technology, e.g. email.

An important issue for the pilots has been those sections of their population that are not engaged with services, whose preferences are not reflected in existing data sets and who cannot benefit in any way if not part of the process.

This led to the development of new and creative ways to consult with communities in order to reach as broad a constituency as possible. This included identifying and contacting a range of groups e.g. community associations, residents and tenants groups, faith groups and other local community organisations.
organisations, as well as patient and carer groups. Individuals could then be involved in one to one interviews, group events, or surveys.

All these approaches benefited from careful preparation, and were followed up afterwards so that patients and service users could learn how their input was used.

**Pilots’ Experience**

**North of Tyne**

North of Tyne organised four patient focus groups in areas where clusters of general practices coalesced. Their existing communications and graphics staff prepared materials for posters, leaflets and local press advertisements. The events focused on:

- Describing the project, its aims and implications.
- The changes service users could expect.
- Seeking users’ views of gaps in services.
- Seeking users’ views on preferences for how these gaps might be filled.

**Tower Hamlets**

In Tower Hamlets a user engagement strategy was prepared which identified three key areas for consultations:

- The concept of self care.
- Patients’ experiences of currently commissioned services.
- What other services people with diabetes felt would help them care for their condition.

A PCT-wide event was held and practices were also supported to set up practice level patient participation groups. Practices paired up to run these and prepared suitable information for patients, taking into account language and literacy issues. Front line staff engaged patients and refreshments were provided to encourage attendance which was excellent, with 80 – 100 attendees. Recruitment via personalised letters from practices proved a much more effective method than advertising through local media.

Following this, Tower Hamlets identified a list of local services and facilities to support self care and sent it to practices for validation and additional content. As a result, a directory was produced containing a menu of options to support patients with self care.

**Calderdale and Kirklees**

Calderdale and Kirklees ran workshops and focus groups for service users in six localities. Chairman, facilitator and speaker notes were prepared to ensure consistency across each locality. Workshops introduced the topic of the Year of Care and invited service users and carers to discuss five key topics related to their diabetes care:

- Information and Education.
- Access to Services.
- Emotional Support.
- Lifestyle/Behaviour Change.
- Social Aspects.

The preferences expressed by service users were fed into their work on provider development (see figure 13).

**User Involvement**

- National Diabetes Support Team (2007) User involvement and diabetes services: Involving voices that are not normally heard (22).
The Year of Care programme has an external evaluation arm. This has two components:

- Evaluating the Year of Care programme including set up and delivery phases.
- Seeking out monitoring tools which could be used long term by clinicians and providers as Year of Care becomes embedded in routine practice.

New Year of Care communities will probably have similar needs and a core set of instruments is available for sharing. This is important as collecting information routinely on the aspects which patients find important, as well as the service finds easy to collect, is challenging. More experience about how to do this will emerge from the next phase of the Year of Care programme.

In the first year the project team might need to establish the standard data that will be gathered. This could include:

- Number of practices involved.
- Number and type of staff trained.
- Number of people involved in care planning.
- List of services identified as forming the menu of options.
- Uptake of services on the menu of options.
- Feedback from care planning consultations into the commissioning cycle.
- Use of templates.
- Inappropriate Accident & Emergency admissions.

The evaluation report for the first phase of the Year of Care, implementation, has been completed. Please contact yearofcare@diabetes.org.uk for more information.

**Pilots’ Experience**

The pilots are finalising their local methodologies for evaluating the Year of Care. This includes measuring the following:

- Cost of services received.
- Consultation experience.
- Service experience.
- Service satisfaction.
- Service capacity analysis.
- Secondary care activity.
- Biomedical data analysis.

Tower Hamlets’ practices have set up focus groups to obtain views from members of their diverse communities. For further information about the evaluation process and results, please contact yearofcare@diabetes.org.uk.

2. Picker Institute Europe (2007) Is the NHS becoming more patient centred?


5. Department of Health (2008) Raising the Profile of Long Term Conditions Care: A compendium of information


7. Matrix Research and Consultancy (2005) Good care planning for people with long term conditions


22. National Diabetes Support Team (2007) User involvement and diabetes services: Involving voices that are not normally heard

APPENDIX I

The Year of Care and care planning demonstration package
The following Year of Care and care planning demonstration package is designed to show an example of the training that has been delivered at the Year of Care pilot sites to meet national quality criteria (16). It is split into the following sections:

1. Programme philosophy.
2. Programme principles.
3. Programme outcomes.
4. Curriculum overview.
5. Programme resources.
6. Programme evaluation.

The final section, Quality Assurance, is still in development: When finished, it will be available at http://www.diabetes.nhs.uk/work-areas/year-of-care.

1. Programme Philosophy

<table>
<thead>
<tr>
<th>About Diabetes and care planning</th>
</tr>
</thead>
<tbody>
<tr>
<td>• People with diabetes are in charge of their own lives and self-management of their diabetes, and are the primary decision makers about the actions they take in relation to their diabetes management*.</td>
</tr>
<tr>
<td>• People are much more likely to undertake action in relation to the decisions they make themselves than decisions that are made for them.</td>
</tr>
<tr>
<td>• People with diabetes need information and both technical and emotional support to make their best decisions about their care.</td>
</tr>
<tr>
<td>• People with diabetes as well as staff, need to be orientated to the care planning approach and what to expect from the experience.</td>
</tr>
<tr>
<td>• Care planning consultations are a meeting of equals, and experts.</td>
</tr>
<tr>
<td>• Care planning needs to be available to all people with diabetes.</td>
</tr>
<tr>
<td>• It is the responsibility of the practitioner to reflect on the style of their consultation and assess how it is supporting the person with diabetes.</td>
</tr>
</tbody>
</table>

* with few and very specific exceptions for example young children, some mental health illnesses and impairments, those whose well being and health decisions are legally made for them by others, temporary physical inability (e.g. following medical treatment, accidents etc).

<table>
<thead>
<tr>
<th>About learning</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Learning about the care planning approach needs to be provided using methods which model the care planning approach.</td>
</tr>
</tbody>
</table>
2. Programme Principles

• The programme is based on evidence-based practice in relation to diabetes and learning.
• The programme is for all staff involved in care planning and all aspects are multi disciplinary.
• People with diabetes are involved in the planning, preparation and facilitation of the programme*.
• The programme consists of:
  o preparation in practice.
  o one day of facilitated learning among peers.
  o time for applying new learning in practice.
  o a half – day of facilitated reflection among peers.
• The programme models the care planning approach, including demonstration and practice of care planning skills.
• Evaluation methods are aligned with the philosophy, learning outcomes and curriculum.
• The programme is audited and quality assured and itself contributes to quality assurance of Year of Care, care planning in a local area / health economy.
• Those facilitating the programme have first attended the programme and reflected on their learning and development as programme facilitators.

*This can be achieved in a variety of ways, including in person, via correspondence, on film and tape and in relation to audit and quality assurance.

3. Programme outcomes

The programme will enable participants to:

• Understand:
  o the concept of Year of Care.
  o the process of care planning.
  o the underlying beliefs and assumptions of the Year of Care project and the care planning approach.
  o the systems that permit effective care planning and allow people with diabetes to be more involved in their care.
  o the difference between a care plan (noun) and care planning (verb).
• Be able to describe:
  o The principles of the care planning approach.
  o Where the Year of Care project fits into the wider field of care for people with long-term conditions.
  o The core competencies and skills required in a care planning consultation.
  o The style, attitude and behaviours that are consistent with a care planning consultation.
  o The implications of care planning from the point of view of someone with diabetes and a healthcare professional.
• Be able to demonstrate awareness of the care planning consultation framework and the competencies and skills required to participate in an effective care planning consultation.
• Be able to construct and explore the Year of Care ‘house’ model in relation to their own beliefs and practice, and place of work.
• Be able to identify potential areas for change relating to:
  o Their own personal attitudes to self management and collaboration.
  o Their behaviours in consultations.
  o Team working, structure and roles.
  o Service structures and systems.
• Have identified goals and developed action plans for:
  o Their own personal development in terms of their consulting style and skills.
  o The changes required in their teams and service structures in order to implement a care planning approach.
• Be able to demonstrate the development of consultation skills appropriate to effective care planning.
• Be able to produce evidence of having reflected on the care planning process.
4. Curriculum overview

<table>
<thead>
<tr>
<th>Preparation in practice (over at least 2 weeks prior to attendance)</th>
<th>Facilitated learning (one day)</th>
<th>Facilitated reflection (half day)</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Self-assessment of personal practice and diabetes related systems and processes</td>
<td>• Introduction, orientation and preparation of environment for learning</td>
<td>• Introduction, orientation and preparation of environment for learning</td>
</tr>
<tr>
<td>• Self-assessment of personal diabetes and practice beliefs and attitudes</td>
<td>• Initial sharing of preparatory assessments</td>
<td>• Structured reflection on implementation of action plan</td>
</tr>
<tr>
<td>• Assessment of enablers and barriers to implementing care planning approach in practice</td>
<td>• Sharing knowledge and defining care planning</td>
<td>• Identification of insights and learning arising from reflection</td>
</tr>
<tr>
<td></td>
<td>• Context clarification</td>
<td>• Sources of ongoing support, QA and audit methods</td>
</tr>
<tr>
<td></td>
<td>• Activities including: house building; identification of roles of those involved; using IT support systems; skills identification, demonstration and practice; goal setting and action planning</td>
<td>• Evaluation and reflection</td>
</tr>
</tbody>
</table>
5. Programme resources

**Essential**

- Preparatory self – assessment and assessment materials
- Care planning consultation demonstration tool/s (script/DVD/scenarios)
- IT template demonstration materials
- Personal goal setting and action planning materials for participants
- PowerPoint slide set of context information
- ‘House’ models for participants (ideally magnetic or jigsaw like kits for personal house-building)
- Evaluation and quality assurance measures
- Personally meaningful take-away materials (e.g. evidence references)
- Structured reflection materials for facilitated half day

**Desirable / To be developed**

- Alternative delivery methods e.g. online, CD-ROM, traditional distance delivery (all with local support from facilitators)
6. Programme Evaluation / Audit Measures

Evaluation of the curriculum – example measures
- Who does the majority of the talking?
- Is an environment created which is conducive to learning?
- Are the learning outcomes covered, and has it been demonstrated that these are understood?
- Does the delivery of the training mirror the care planning approach?
- Does the training generate, and answer, questions?
- Do participants arrive already stimulated by the topic?

Evaluation / Audit of the Programme – example measures
- Delivery of the learning outcomes.
- Change in the way in which care is delivered, assessed against improvements in scores in relevant evaluation resources, for example the Consultation Quality Index-DM (CQI-DM), changes in systems and processes to facilitate the care planning approach and audit of care planning components.

7. Programme Quality Assurance (QA)

A QA tool for commissioners is in development. It will include:

1. A checklist for commissioners to employ to assess whether the training they have commissioned consistently meets the specification of Year of Care training, including philosophy, learning outcomes, content and resources.

2. A measure or measures to be completed by participants to assess the learning experience and the provision of all parts of the programme as described. QA needs to be undertaken internally and externally.

Both QA and audit awareness activities must be included in the content of preparation of learning facilitators for care planning, which may be undertaken following participation in this programme. Audit and quality assurance will be key evaluation markers of the programme.
APPENDIX II

NHS Yorkshire and the Humber IT templates

IT templates capable of capturing care planning data have been designed in TPP SystmOne by NHS Yorkshire and the Humber. TPP SystmOne and NHS Yorkshire and the Humber are happy for this work to be shared on condition that the following principles are upheld:

1. This information is to be used for the benefit of patient care.
2. This information should not be used by others to make commercial gain from replicating the approach in their own systems.
3. The joint work of NHS Yorkshire and the Humber and the Year of Care teams in producing these initial templates is recognised.

The key elements of the IT templates developed are that, in addition to biomedical data, they capture:

1. The patient’s story/perspective.
2. The domains of care (i.e. knowledge and health beliefs; emotional; behavioural; social; and clinical).
3. The patient’s goals.
4. The actions to be taken to help the patient achieve their goals.
5. The outcomes.

It is critical that systems are capable of recording this data as well as providing the essential administrative tasks required for care planning (e.g. sending out letters to patients). Below is a screenshot of one of the care planning templates.

If you have any queries relating to these templates, the principles described above or any other aspect of care planning data capture, please contact yearofcare@diabetes.org.uk in the first instance.
For further information on Year of Care contact:

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Diabetes UK

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Email: James.Thomas@diabetes.org.uk

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