How can I help reduce healthcare associated infections?

Infection control is important to the well-being of our patients and for that reason we have infection control procedures in place. Keeping your hands clean is an effective way of preventing the spread of infections. We ask that you, and anyone visiting you, use the hand sanitiser available at the entrance to every ward before coming in to or after leaving the ward. In some situations hands may need to be washed at the sink using soap and water rather than using the hand sanitiser. Staff will let you know if this is the case.

Contact us:
Tel: 01494 426976 or 01296 566045
Email: buc-tr.cypphysio@nhs.net
website: www.buckshealthcare.nhs.uk/CYP

Chronic Fatigue Syndrome in children & young people

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What happens next?

The attached flow chart shows the process of action you can expect once a referral has been sent by the Paediatrician to the Community CFS team:

- Questionnaire sent to family for completion and return to the CFS team prior to an appointment being offered.
- Initial appointment with a Physiotherapist and/or Occupational Therapist involves:
  - Explanation of process
  - In-depth review of current functioning
  - 2 week activity diary given for completion
- The second appointment involves:
  - Discussion of activity diary
  - Joint setting of short/long term goals with young person and their family
  - Care plan agreed, including involvement of other team members
- Visits arranged by ‘Key worker’ to review progress and to set further goals.

Continued liaison and review of care plan with other services as necessary.

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**Introduction:**
You have been referred to the Children and Young People’s Community Chronic Fatigue Syndrome (CFS) Team for assistance with your symptoms. This leaflet is to inform you about who we are and what we do.

**Understanding Chronic Fatigue Syndrome:**
CFS is a condition that has fatigue or tiredness as one of the main symptoms. Other associated symptoms may be joint pain, headaches, sleep disturbance and mood changes.

A specific cause of CFS is not yet fully understood. However, some patients have noticed an onset of symptoms following a viral infection or prolonged adrenalin release due to sustained periods of mental, emotional or physical stress.

When a person has been unwell for a long time and has been unable to carry out normal activities, his/her body can become deconditioned making CF symptoms and mood feel worse.

The CFS Team for Young People:
The professionals who comprise the Community Paediatric Team include Paediatricians, Physiotherapists, Occupational Therapists, Psychologists, School Nurses and Home tutors. Occasionally a Dietician, may also be involved. The people that you will meet will be dependent on your individual needs.

The team members liaise regularly and can become involved at any stage. We will also keep your GP informed.

The primary aim of the Community Paediatric Team is to regain a lifestyle that is acceptable to for you and your family.

**Deciding the best way forward:**
Following your diagnosis of CFS your Paediatrician will refer you to The Team. You will then be sent a questionnaire to complete which will provide us with more specific information on how your lifestyle and everyday activities are being affected.

Following this, you will then meet representatives of The Team to gain a more in-depth picture of your current level of sickness, activity and coping skills. They will take this information to the rest of The Team and discuss what would be the best way forward for your care and provide a suggested plan.

**Planning Intervention:**
Each young person will have his/her own Key Worker, who will discuss and plan the best way forward with you personally.

Support can be given on any, or all of the following:-

- Advice and programmes on how to manage energy and activity levels.
- How to deal with feeling scared, i.e. anxious, or feeling low.
- Getting back into school and social activities.
- Decisions about medicine which might help with pain or sleep should you need it.

This support will be co-ordinated between Health, Education and Psychology Services.

**Your Progress:**
Your Key Worker will be your main point of contact for ease of communication. They will most typically either be a Physiotherapist or Occupational Therapist.

Your Key Worker will review your progress with you, and The Team, at regular intervals to ensure that effective input is being provided.

**School**
We recognise that school is an important part in a young person’s life, not only academically but socially too.

We also recognise that when a child has been out of school for a period of time, returning back can be daunting. Therefore, good communication between the CFS team, the school and the young person is essential.

If you have any questions about this leaflet please do raise them at your assessment.