

Communication can be a difficult process *with ICU patients*. Patience is required by all involved and it is common to use a variety of methods. Please ask staff on the Unit for help and guidance.

Sometimes, patients can be confused and forgetful, and it is necessary to repeat information for them again and again. (Please see Hallucinations leaflet or ask a staff member for more information.)

### Points that are common from patient feedback are:

- Where am I?
- Why am I here?
- Why can't I talk?
- Why can't I eat and drink?
- What are all the alarms for?
- What are the tubes for?
- I'm uncomfortable
- I want to watch TV/listen to music etc.
- What time is it?
- When will I get better?
- I want to go home
- Why are you leaving me here? Why can't I go with you?

If you are concerned about communicating with your relative, please tell a member of staff or contact the Rehab after Critical Illness team (details on the back page).

## Rehabilitation after Critical Illness

Level 3 South Block

T: 0118 322 7248

E: [RaCI@royalberkshire.nhs.uk](mailto:RaCI@royalberkshire.nhs.uk)

The Rehab after Critical Illness team can be contacted on the above number if you have any concerns regarding your relative's altered mental state. (A message can be left if it is out of office hours.)

The team will visit the patient when they go the ward as part of the normal rehabilitation process for continued observation if there are any concerns.

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ICU support webpage –  
[www.readingicusupport.co.uk](http://www.readingicusupport.co.uk)

Royal Berkshire NHS Foundation Trust  
London Road  
Reading RG1 5AN  
Telephone 0118 322 5111  
[www.royalberkshire.nhs.uk](http://www.royalberkshire.nhs.uk)

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**NHS**

Royal Berkshire  
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# Voiceless – why can't the patient talk?

Information for patients,  
relatives and carers

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Some patients in ICU cannot talk mainly due to having a 'breathing tube' in place.

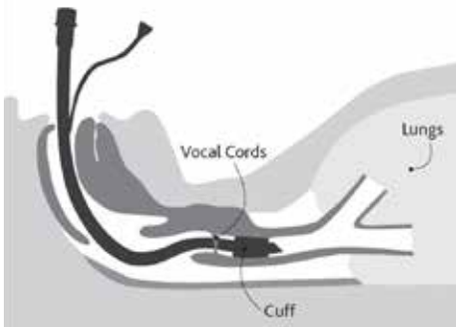
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The patient will be on a ventilator ('breathing machine'). This might be through a tube in the mouth or in the throat.

Both of these tubes go into the windpipe and past the vocal cords. Each of these tubes has a 'cuff'. When it is inflated, this helps the breathing machine deliver all the support needed and helps keep the tube in place. When the cuff is inflated, no air can pass up to the vocal folds. It is the air flowing over the vocal cords that helps the vocal cords vibrate, creating the sound which is your voice.

Cuff up = No voice

It is not possible for the patient to drink or eat at this time either as the cuff affects the ability to swallow.



The patient might at this time believe that they can speak, and from patient experience, believe they can hear themselves talk.

### Patient Experience

*"Having been on a ventilator for a long period of time through a tracheostomy the frustration and fear you feel at not being able to speak, is overwhelming. I was not aware at the time I was attached to a ventilator; you can't feel it and you can't see it. I believed I was speaking quite clearly to people, I even believed I could hear my own words. It is hard to describe the feeling of loneliness you get when you desperately want someone to understand you. Only afterwards do you also realise the frustration others feel when they want nothing more than to understand you. Despite the frustrations, I never forget, the ventilator kept me alive. Seeing a simple drawing which was done for me of how a tracheostomy prevents you from speaking made such a difference to me, I felt relieved and educated that I now knew why".*

*Jason Boyes*

Once the patient is well enough not to need the breathing machine all the time, and if the patient has a tracheostomy, the cuff can be deflated. If the patient is not "distressed" by this (e.g. finding breathing more difficult, oxygen levels falling, coughing continuously) a 'speaking valve' can be tried on the end of a tracheostomy. The healthcare professionals will decide when this is appropriate. The speaking valve is not guaranteed to work!! Often, patients do not manage this as they might be too weak.

The patient will not speak with their normal voice even with the speaking valve in place. When a patient is on a breathing machine, they might be "suctioned" – that is secretions removed from the chest that they cannot always clear themselves. Feedback from patients is that being told why this is being done is important to them, as it can be an uncomfortable procedure and this explanation needs to be given more than once.

### Other ways of communicating

- 1) The patient might mouth words. This may be a very frustrating time for all involved. Whilst this works for some people, lip-reading can be very difficult.
- 2) The patient might try to write for you. Some assessment of the patient is involved before attempting this as they might not have the strength and coordination. Again, the patient might believe what they have written is clear and makes sense.
- 3) Alphabet/picture boards – the patient uses their fingers or eye gaze to point to a picture or a letter to spell a word.
- 4) We have other high-tech aids to help with communication. We have available iPads and a computer with pictures and symbols that can 'speak' out loud typed words or selected pictures. Our computer has switch access and electronic eye-gaze technology.