OPINION

Chronic cough: from a complex dysfunction of the neurological circuit to the production of persistent cough

Adalberto Pacheco

ABSTRACT

Chronic cough or cough that lasts more than 8 weeks, once a chest x-ray and spirometry are confirmed normal, is caused by an alteration in a section of the route between peripheral receptors, mainly in the upper and lower airway and oesophagus, spinal cord and the cough centre in the brain stem involving the cortex. These mechanisms of cough have their homology in the circuit of chronic pain, and on that basis, should expand future research of chronic cough. Clinically chronic cough is easy to diagnose by an excessive response or hypertussia to low-intensity stimuli or banal stimuli, which we now call hypersensitivity cough syndrome, quantified by a positive reflex cough with capsaicin or citric acid. However, hypersensitivity cough syndrome can be impossible to quantify in the laboratory when the hyper-responsiveness originates in the central nervous system. This is normally caused by excessive peripheral input or convergence of stimuli from different sources. Once central hypersensitivity is acquired, peripheral input is not as important for activation of the cough.

The prevalence of chronic cough (CC) reported in studies in Europe and the USA varies between 9% and 30%.1 However, in two-thirds of patients the cough is completely or partially resistant to conventional treatment.2 In the last decade, the recognition of the similarities between the neurobiological mechanisms underpinning cough and pain has been an important development.3 To date, chronic pain has been studied in considerable depth, and now it is time to apply the findings from chronic pain studies to the management of CC.

The mechanism of cough is structured as a neurological circuit connecting the periphery to the central nervous system (CNS). In some instances the cerebral cortex is incorporated into the circuit in the form of the sensation of urge to cough, or voluntary cough, as opposed to the other kind of cough, the cough reflex. As in chronic pain, in CC the clinical problem arises when there is a prolonged imbalance between stimulus and response which impairs patients’ quality of life and causes them to cough in response to mild stimuli, or even in response to innocuous stimuli such as talking or laughing. For this reason, this prolonged cough is now catalogued as a Cough Hypersensitivity Syndrome4 or sensory hyper-responsiveness to numerous mechanical or chemical stimuli occurring at different anatomical sites, especially the upper and lower airways and the oesophagus. These stimuli are transduced into electrical action potentials in the terminal fibres of the vagal or trigeminal neurons, all of which have their cell bodies in the spinal cord. These fibres, which are involved in the cough circuit, have been studied in depth in laboratory animals. Basically, there are three types5: inflation and deflation receptors—rapidly adapting receptors and slowly adapting receptors—which do not play a major role in the production of cough; myelinated Aδ fibres (termed ‘cough receptors’ by some authors) which respond to mechanical stimuli and acids; and unmyelinated C-fibres, which respond to thermal and chemical irritants such as capsaicin.

The first mechanism in the production of cough is the increased excitability of peripheral nerve endings to a given stimulus. Many types of chemical stimuli can elicit a neuroinflammatory response; these stimuli have been referred to collectively as an ‘inflammatory soup’.6 The activation of TRPV1 and TRPA1 receptors present in C-fibres in the respiratory system has been explored in depth. One study in humans found that prostaglandin E2 inhalation increases the cough response to subsequently inhaled capsaicin, but the activation of Aδ fibres may also be important; in a guinea pig model, after allergen-induced mast cell activation, a state of hypersensitivity to the mechanical stimuli that activate these nociceptors was observed.7

If the increase in excitability occurs in two different types of terminal fibres, this may lead to the phenomenon of convergence and synergy prior to cough.8 Confirmation of this kind of convergence was found when either bronchoconstriction (rapidly adapting receptor-mediated) or acid (C-fibre-mediated, in the distal third of the oesophagus) was provoked in guinea pigs; it was shown that neither event was sufficient to trigger cough alone, but simultaneous stimulation was able to cause coughing.9 In humans, oesophageal acid perfusion caused cough only in subjects with prior states of airway irritation such as asthma, but not in healthy subjects.10 This conjunction of two sites of activation has been identified by some authors as the origin of visceral hypersensitivity, a useful concept to explain that even normal levels of acid in the distal third of the oesophagus in healthy individuals may, in the case of concomitant CC, increase cough sensitivity.11 A similar convergence mechanism has been demonstrated in the nasal cavities and trachea to explain CC originating in the upper respiratory tract.12 These results suggest that action on two or more potential sources of cough

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such as the oesophagus and upper or lower airways may be beneficial for treatment of CC. This particular relationship was targeted in a clinical study of the connection between gastrooesophageal reflux and eosinophilic airway inflammation.13

The second mechanism involves more active participation on the part of the CNS, in the form of central sensitisation. Repeated peripheral stimuli can create a state of central hypersensitivity that may be reversible, although if the activity of nociceptors persists over time it may trigger a hypersensitivity known as spinal long-term potentiation.14 There are two interconnected ways in which this link between the periphery and the CNS can develop. One is via the interaction between terminal fibres, that is, when sensory input carried by the C-fibres to the cough centre (the nucleus tractus solitarius located in the brainstem) activate other nerve endings, whose sensitivity to these stimuli is probably increased by the synaptic plasticity in the cough circuit15; this has a synergistic effect, and produces cough.9 In one animal study, after stimulation of C-fibres at sites where there are no Aδ fibres, such as the periphery of the lung or the oesophageal mucosa, an increase in input from Aδ fibres in the mucosa of the larynx, trachea and bronchi was observed; then, the joint action of the Aδ fibres and the C-fibres produced an increase in central sensitivity.9 In the context of chronic pain, an identical phenomenon occurs known as secondary hyperalgesia;16 in cough, the condition is known as secondary hypertussivity, which refers to the way sensitisation originating in a specific somatic area is perceived as being more extended than it really is; for example, a peripheral stimulation of acid in the oesophagus might be perceived by the CNS as being caused by receptors at other sites such as the larynx or the airway.

The second potential influence of peripheral stimuli on the CNS is the effect exerted on the first neuron of the cough circuit by the microglia, a set of bone marrow derived cells that participate in primary immunity and are activated in chronic circuit by the microglia, a set of bone marrow derived cells that may be that its modulation in the spinal cord in response to cough stimuli.17 A last factor to bear in mind is the descending control on the nucleus tractus solitarius (stimulatory and inhibitory), as in the pathophysiology of chronic pain; in turn, this control is influenced by the synapses in the pathway through the periaqueductal grey and the limbic brain, with the result that anxiety and depression may modulate the final efferent motor impulse.3

Figure 1. Feedback neurological circuit of chronic cough.
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