Carcinoma of the lung: warts and all

A F Markham

It is always a salutary experience to be confronted by one's own ignorance. The consequences, however, can often be worthwhile. In my own case, the paper by Soini et al on pp 887–93 of this issue of Thorax represents a good example of this. The issues it raises should certainly provide food for thought for a wide range of respiratory physicians, other clinicians, pathologists, and medical scientists.

Firstly, I should confess to the many misconceptions under which I first read the paper by Soini et al. I have followed the debate and controversies surrounding the role of human papilloma viruses (HPVs) in squamous cell carcinoma of the cervix for the last decade at a very superficial level. Initially, much of that controversy was fuelled by technical difficulties in reproducibly detecting low levels of viral DNA in clinical specimens during the era before polymerase chain reaction (PCR). Even when these practical problems were surmounted, the causative role of HPV and, in particular, the high risk strains HPV 16 and 18, remained controversial because they were almost certainly not present in all squamous cell cancers. The current consensus seems to be that HPV16/18 (or sometimes 31/33) are present in about half of squamous cell carcinomas of the cervix and probably initiate transformation to malignancy in these cases; in other cases the disease may well have arisen secondary to a sporadic mutation event in the p53 gene without the involvement of HPV. The plausibility of this dual mechanism model is strengthened by the fact that both impact directly on p53 protein function. I imagine that this might be the level of understanding of this interesting model of malignancy amongst most respiratory physicians. I was vaguely aware that HPV16/18 had been implicated in the development of squamous cell carcinoma of the oropharynx, upper oesophagus, and anus. This was consistent with the presence of squamous epithelium at those sites. I therefore assumed that the Finnish group's work describing the presence of HPV in squamous cell carcinoma (and other cancers) of the lung was simply another example of a well recognised phenomenon and therefore not of particularly great novelty or importance.

To my surprise, there have in fact been relatively few previous studies of possible involvement of HPV in lung cancer over the past 10 years reported in the world literature. The scarcity of these observations has reduced their impact. The limited published data suggest that HPV is not infrequently present. Given that squamous cell carcinoma represents some 40% of this most common of human malignancies, these small numbers of observations assume a considerable significance. The findings of Soini et al and others are remarkably consistent with those made by many workers in the cervical cancer field. To a first approximation, bronchial squamous cell carcinoma and other histological types of lung cancer display either the presence of oncogenic HPV species or spontaneous p53 mutation. The fact that Soini et al also found HPV in some lung adenocarcinomas probably need not concern us too greatly at this stage. The complex mixed cellularity in many lung cancers is well recognised, and the widespread occurrence of p53 mutation in lung cancer is well known. The fact that this malignancy is very strongly associated with cigarette smoking is thus hardly surprising. Marinating bronchial epithelial cells which have been infected with a transforming strain of HPV, or have undergone a p53 mutation, in the mixture of carcinogens provided by cigarette smoke, has to be a recipe for disaster.

It is worth digressing briefly to consider the mechanisms by which HPV16/18 are thought to cause transformation of cells. These viruses encode transforming proteins, E6 and E7, which are thought to exert their respective effects by interfering with the function of p53 and Rb tumour suppressor proteins directly. E7 binds directly with Rb and prevents its function as a regulator of the cell cycle. The action of the E6 proteins is a little more complex. In the HPV infected cell E6 interacts with a human protein, the E6-associated protein (E6-AP), and E6 and E6-AP are thought to act together as a ubiquitin ligase which recruits a human cellular ubiquitin conjugating enzyme to ubiquitinate p53. The actual mechanism may involve ubiquitination of the E6/E6-AP complex which itself transfers the ubiquitin to p53. Ubiquitination of a cellular protein targets it for proteolysis. Thus, in a cell infected with HPV, wild type p53 will be rapidly and extensively degraded with the result that the cell effectively has a p53 null phenotype.

What draws the two putative aetiologies of squamous cell carcinomas together very clearly. Initial cellular transformation by virtue of loss of p53 function may occur either by mutation in the p53 gene itself or by destruction of the p53 protein secondary to HPV infection. The fact that some squamous cell carcinomas have both mutant p53 and HPV infection is explicable by the resistance of some p53 mutants to E6 driven proteolysis. All these events are now fairly well understood at the molecular level. The functions of p53 and E6 have been widely studied, the human E6-AP gene has been cloned and expressed to provide protein reagents for experimentation, and several groups are currently identifying ubiquitin conjugating enzymes (UBCs) which target p53. Indeed, the first experiments have been reported in which inhibition of E6-stimulated p53 degradation has been achieved. This has obvious potential therapeutic possibilities in that treated cells would regain normal p53 function which might reasonably be expected to suppress the malignant phenotype.

What happens next? Obviously it is now essential that the data of Soini et al and their predecessors in the literature are confirmed by others. The high frequency of this disease should mean that this phase is not protracted. Technically this will not be a trivial exercise given the need to detect very low levels of infection by in situ hybridisation or the possibility that minor sequence variation in HPV isolates might cause PCR failures. However, on the assumption that the data are reproducible (and there always tends to be controversy in these exercises because of false negatives), a number of other issues arises. The bronchial epithelium is not classically characterised as squamous. What then is the cell initially infected with HPV? The possibility that these cells are shed from the squamous epithelium of the oropharynx is less convincing because squamous cell carcinoma is found both in the lung peripheries and upper lobes. However, there is a recognised association between oral papillomatosis and squamous cell lung cancer. Hopefully, the new observa-
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squamoucell carcinoma of the oropharynx, upper oesophagus, and anogenital regions did not represent sufficient numbers of patients to justify the levels of investment the pharmaceutical industry would need to make. Inclusion of bronchial squamous cell carcinoma in the list of HPV based cancers would change these financial considerations completely. The p53 ubiquitination mechanism provides a possible starting point for pharmacological intervention. Ironically, this may provide benefits to another small group of patients which suffers the ravages of HPV. Renal transplant patients, presumably as a result of their immunosuppression, have dreadful problems with papillomatosis and, in particular, patients with 10 year graft survival have a 30% incidence of squamous cell carcinoma of the skin, as well as considerable risk of lesions in the anogenital tract. Perhaps they will also be the indirect beneficiaries of the intriguing observations of Soini et al.

I have confessed to my own ignorance. With luck the pharmaceutical industry will not be using the same excuse for very much longer. Recent description of a recombinant vaccinia virus encoding papillomavirus E6 and E7 and its use for immunotherapy (in cervical cancer) is particularly timely and exciting. Furthermore, given that three of the four main “Health of the Nation” national targets for cancers are a 30% reduction in lung cancer deaths in men (15% in women) by 2010, a 20% reduction in the incidence of invasive cervical cancer by the year 2000, and a halt in year-on-year increases in the incidence of skin cancer by 2005, perhaps our masters in the R & D function of the NHS should also be taking the study of HPV rather more seriously.

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Molecular Medicine Unit, University of Leeds, Clinical Sciences Building, St James’s University Hospital, Leeds LS9 7TF, UK

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