RESEARCH UPDATE

Pulmonary metastasectomy in colorectal cancer: the PulMiCC trial

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ABSTRACT

PulMiCC (Pulmonary Metastasectomy in Colorectal Cancer) is a randomised controlled trial funded by Cancer Research UK. Patients with a history of resected colorectal cancer who are found to have pulmonary metastases are first registered for evaluation and, if subsequently eligible for the trial, are invited to be randomly allocated to ‘active monitoring’ or ‘active monitoring with pulmonary metastasectomy’. The clinical outcomes are overall survival, relapse-free survival, lung function and patient-reported quality of life.

Pulmonary metastasectomy—that is, the surgical removal of nodules of metastatic cancer from the lungs—is very well established and widely accepted practice, most commonly for colorectal cancer. From that starting point, the Pulmonary Metastasectomy in Colorectal Cancer (PulMiCC) trial is best understood as an investigation to discover if active monitoring without pulmonary metastasectomy might result in survival similar to that reported in surgical series when like is compared with like. The evidence available for the effectiveness of metastasectomy is in the form of retrospective case reviews and observational studies. These have been collated in a formally conducted systematic review and a quantitative synthesis of data from 3504 patients in 51 studies. There were no randomised trials. The 5-year survival rates of patients who had pulmonary metastasectomy are of the order of 40%. In only two reports is a denominator provided. These show that metastasectomy was performed in a selected 1–3% of patients with recurrent cancer. Thirty years ago the suggestion was made that this better than expected survival might be an effect of selection rather than being predictive of the surgical effectiveness of pulmonary metastasectomy.

However, practice has gone well outside the bounds set in the observational studies. In a recent survey among members of the European Society of Thoracic Surgeons (ESTS), multiple metastases were not a contraindication for 85% of responding surgeons, an interval of less than a year since the primary operation was acceptable for 64% and synchronous metastases for 50%. Whether the reported good survival rates are due to surgery or selection, it cannot be assumed that they will be replicated in a practice which has strayed well outside the criteria employed in the selection of the 3504 patients in the quantitative synthesis. In contrast to the stated practice of the members at large, the ESTS Working Group was unable to offer recommendations for pulmonary metastasectomy on the basis of its findings.

The design of PulMiCC is founded upon an uncertainty that must exist in the advice given to many of these patients. Consider two scenarios where advice for or against metastasectomy is reasonably consistent. Patients who have a cluster of the favourable features, such as a single pulmonary metastasis which is evident only after several years and is growing slowly, might be offered metastasectomy as part of widely accepted current practice (although it should be noted that effectiveness of metastasectomy has not been shown and these patients are eligible for randomisation within PulMiCC). In contrast, patients with multiple bilateral metastases, evident within months and growing, are generally advised against metastasectomy. However, between these extremes are patients with a combination of some favourable and some unfavourable features; they are neither an easy ‘yes’ for metastasectomy nor a confident ‘no’. If there is uncertainty, unbiased allocation is as good a basis as any other for making the decision.

PulMiCC has a two-stage consent process (figure 1) derived from that used in the Mesothelioma and Radical Surgery (MARS) trial where it proved an effective strategy to recruit sufficient patients into a very challenging trial. In the first stage of PulMiCC, any patient with one or more pulmonary nodules which might be from colorectal cancer can be invited to enter the study with no commitment to randomisation. DVDs for medical staff and for patients have been prepared based on evidence-based best practice in presenting uncertainty to patients in a way that is open, even handed and, at the same time, supportive. The notion that haste or urgency are helpful should be
dispelled. Even if it is not easy to put these matters to patients, for those with demonstrable stage IV colorectal cancer, time measured in days or a couple of weeks is better spent in careful evaluation than hastening towards a thoracotomy.

During this first stage of PulMiCC the operative and pathological details of the primary cancer are reviewed. Where there is a solitary nodule, the possibility of primary lung cancer should be considered by the lung multidisciplinary team. If not already done, PET/CT is arranged. In some cases percutaneous biopsy will be advised. With regard to chemotherapy, we found no evident consensus. Some oncologists give chemotherapy routinely on discovery of recurrence, some believe it has a place along with patient-reported quality of life outcomes, pulmonary function and survival.

The trial is open for recruitment in an increasing number of UK centres. Full details of PulMiCC can be found on the website (http://www.rbht.nhs.uk/PulMiCC/).

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REFERENCES


**Images in Thorax**

**Pulmonary hamartoma mimicking primary bronchoalveolar cell carcinoma**

A 52-year-old man with transitional cell carcinoma underwent a chest CT because of a newly developed pulmonary nodule observed on a chest radiograph. In addition to the nodule, a 2.8×2.4 cm well-circumscribed, pure ground-glass opacity with air cyst formation was incidentally found (figure 1). There was no evidence of calcification or fat, and primary bronchoalveolar cell carcinoma was highly suspected. When performing preoperative CT-guided needle localisation of the opacity 2 weeks later, we found that the lesion remained unchanged. After video-assisted thoracoscopic wedge resection,

Figure 1  Chest CT image with soft tissue window settings (A) and lung window settings (B) showing a 2.8×2.4 cm well-circumscribed, pure ground-glass opacity with air cyst formation in the right upper lobe. There was no identifiable soft-tissue component, calcification or fat within the lesion.

Figure 2  Gross morphology of the specimen showing a well-circumscribed, spongy and whitish tumour (arrows in A). (B) Pathologic examination demonstrating that the lesion was composed of cartilage, fat and smooth muscle with benign bronchial epithelial cells (H&E staining ×100).