BOOK REVIEWS


Just as Marks is to Spencer, so St Mark’s is to polyposis. The condition may have remained the same but only recently has the term familial adenomatous polyposis become common currency. It has evolved through stages variously as familial polyposis coli, familial multiple polyposis, multiple intestinal polyposis, familial adenomatosis coli, familial polyposis coli – and just about every other combination of the terms – until arriving at its latest form. Indeed, in the way that terminological difficulties are now resolved, a meeting of 30 of the great and good in colonic polyps from 11 countries met at Leeds Castle in June 1985 to resolve the nomenclature. Those Euro-sceptics among the gastroenterological community might be distressed to hear that the horizon of polyposis groups has now extended from a castle in Kent to Euro-FAP, which is supported financially by the European Union.

It is obviously worthy that such experts in a comparatively rare condition should not only get together to share their experiences, data and, above all, their patient bases. It must be recognised, however, that all of the fundamental work in setting up registries, which has proved such a fruitful foundation for research, have come from the effort of three people who worked at St Mark’s Hospital: Lockhart-Mummery, Dukes, and (at that time) Dukes’s junior laboratory technician – Bussey. The tribute that the present volume pays to Bussey is clearly justified by his enormous and meticulous contribution, especially in establishing the St Mark’s Polyposis Register (in 1924 or 1925, there is some uncertainty). How Bussey is made abundantly clear in the preface and generous foreword. The other name that towers throughout this book is that of Gardner, who sadly died some six years ago at the very time that molecular biology was coming in on the gene defect in Gardner’s syndrome. The interesting suggestion is made here that, as a mark of respect and acknowledgment of their contributions, familial adenomatous polyposis might be renamed Bussey-Gardner polyposis. It would certainly be a fitting memorial but I just wonder how easy it would be to eradicate the familial and polyposis from the title of the condition and, of course, although the edges of Gardner’s syndrome have become somewhat blurred in recent years, it does stand as a fairly established entity as St Mark’s Hospital closes its doors and moves to new pastures within Northwick Park Hospital elsewhere in London, I would have thought there was a good case to be made for renaming the condition St Mark’s syndrome.

This is a superb book, which has been lovingly edited by three of the senior surgical staff at St Mark’s Hospital. Their affection for those involved in the history of this condition, as well as their benefit of such an embalmed subject, permeates through every page and surely none can challenge the credentials or the authority of any of the people involved with this monograph. The history, pathology, genetic, and clinical evaluation of patients are covered in separate chapters. All are readable, and the chapter on genetics is particularly clear. As with all the very best of British surgical teaching, anecdote abounds. Indeed, the potential of such an embalmed subject to teaching are not restricted to surgeons: there is an exceptionally witty parody of Lewis Carroll’s ‘Father William’ as applied to polyps and was penned by Cuthbert Dukes. I think this is good enough to share with Gut’s readers.

‘You are old, Father William,’ the young surgeon said, ‘And your colon from polyps is free. Yet most of your sibling are known to be dead – a really bad family tree.’

‘In my youth,’ Father William replied with a grin, ‘I was told that a gene had mutated, That all who carried this dominant gene To polyps and cancer were fate’d.’

‘I sought for advice from a surgical friend, Who sighed and said, “Without a doubt Your only escape from an untimely end Is to have your intestine right out.”’

‘It seemed rather bad luck – I was then but nineteen – “So I went and consulted a quack, Who took a form hold on my dominant gene And promptly mutated it back.”’

‘This,’ said the surgeon, ‘is something quite new And before we can ascribe any merit We must see if the claims of the fellow are true, And observe what your children inherit!’

If this book has a fault, it is that the last 60 odd pages are devoted to ‘other polyposis syndromes’. Inevitably, this does not have the depth of the earlier chapters on FAP, and mentions some of the cancer family syndromes, which are, of course, strictly ‘non-polyposis cancer syndromes’. I think this departure does perhaps illustrate the difficulties of not knowing quite where to stop in the search to be comprehensive.

I cannot believe there is a single practising gastroenterologist who would not benefit from studying this book.

IAN FORGACS

This title may be purchased from the distributor: Bookpoint Limited, 39 Milton Park, Abingdon, Oxford OX14 4TD. Tel: 01235 400403; fax: 01235 821511.