Surgical oncologists need constantly to balance delivering the best long-term cancer-free outcomes against preserving function and facilitating rapid recovery. Minimally invasive surgical techniques and alternative methods of tissue ablation such as high intensity focused ultrasound (HIFU), radio-frequency ablation, cryotherapy and laser-based photodynamic therapies are being tested to address this need. One major question is which men require radical treatment because without it they are at risk of progression, metastasis and death, and which men might be managed by careful monitoring. Another question addressed by this paper is whether some men with low or intermediate risk prostate cancer (PC) might wish to have surgical treatments which have an increased risk of leaving tumour behind, but which have better side effect profiles. The crucial question they raise, however, is whether the trade-off between better quality of life and the risks of leaving some residual local tumour can be safely calibrated. The reason this is important is because evidence from meta-analyses shows that positive margins are associated with increased rates of biochemical recurrence (by 35%–50%),1 in men having standard radical prostatectomy, although in T2 Gleason 6 disease, the increased risk of biochemical recurrence may be modest.

When new techniques are developed, researchers have an obligation to determine that they are safe and effective and that the process of introduction does not harm patients. These challenges are acute in the management of PCs. Sood and colleagues from the Vattikuti Urology Institute at the Henry Ford Hospital in Detroit describe in this issue of the journal a modification of the process of introduction does not harm patients. These challenges are acute in the management of PCs. Sood and colleagues from the Vattikuti Urology Institute at the Henry Ford Hospital in Detroit describe in this issue of the journal a modification of existing techniques of robotic-assisted laparoscopic prostatectomy (RALP) for the treatment of localised PC. Do they pass the test?

Current evidence suggests that mp-MRI and targeted biopsy coupled with systematic biopsy is the most accurate way of identifying men with significant cancers at diagnosis.2–4 But how accurate are such approaches in picking up additional clinically significant cancers (Gleason Group ≥2) within a prostate known already to harbour a cancer? While the precise figure is open to debate, it is clear that men with clinically significant disease frequently harbour multifocal cancers. It is also clear that many of these cancers are missed by current diagnostic approaches and are distant from the ‘index’ cancer.5 6 Furthermore, it is now clear that within the prostate there are significant ‘field effects’ and that while some cancers have evolved from a single clone, others are truly polyclonal in origin.7 For these reasons, reservations continue to exist around the use of focal therapy where only the cancer(s) which can be seen (the index or dominant lesion) are treated. Conversely, the rationale for testing conservative approaches is that Active Surveillance is proven to be a valid approach for Gleason Grade Groups 1 and 2, since most low risk and intermediate risk PCs do not cause symptoms, metastasis and death within a 10-year time frame8 and many low and intermediate risk cancers stay of the same grade over time.9 However, it also clear that dedifferentiation can occur over time in some men.10 11

Progression and death rates in such men at 10 years are extremely low following surgery, radiotherapy and active surveillance8 and biochemical recurrence following surgery in Gleason Groups 1 and 2 is ~20% at 10 years,12 13 so measurement of safety for new approaches can only be defined in the long-term. Nevertheless, increased rates of early cancer recurrence are concerning. Another open biological question is if creating a wound within the prostate either by HIFU or partial prostatectomy, resulting in an environment rich in cytokines and growth factors, might actually promote long-term cancer growth if malignant cells are left behind.14

Menon and colleagues have tested the idea of carrying out a standard robotic-assisted
nerve-sparing radical prostatectomy on the side of the main lesion and carrying out a fascial-preserving approach on the contralateral side in a selected group of men who might have been suitable for HIFU/focal therapy. The purpose was to remove all the prostate and all the cancer on the side of the dominant lesion; on the other side, the fascia and a ‘thin rim of prostate’ (noted as 5mm in one part of the manuscript and 5–10mm in another) were deliberately left behind. They termed this partial removal as ‘precision prostatectomy’.

A preliminary study of 100 radical prostatectomy samples was carried out, modelling the presence of cancer in the rim that would have been likely to be preserved with ‘precision prostatectomy’ and therefore what cancer would have been left behind after the new operation. This exercise suggested that 35% would have had cancer left behind, 14% having clinically significant disease. However, only 25 of these 100 cases would have been suitable for HIFU or focal therapy, which was one inclusion criterion for the ensuing pilot clinical study of eight men with good sexual function (four with Gleason 3+3; four with 3+4 disease). It is not clear from the paper how many of these 25 men would have had cancer left behind if they had undergone a ‘precision prostatectomy’. It was surprising to me that routine postoperative mpMRI and modern imaging with PSMA PET-CT was not performed or reported.

At 12 months, the functional results were excellent with all men having good sexual function and continence. This compares with the authors’ previous potency rates of ~80% following standard RALP (although ~50% of the 80% used PGE-5 inhibitors). As far as cancer outcomes are concerned, on the side of the ‘precision prostatectomy’ positive margins were found in three men (37.5 %) including two of the four men with Gleason 3+4=7 disease. These numbers are small, but in these men with ≤T2 disease, they translate into positive margins in 50% of men with Gleason 3+4=7 disease and 25% of men with Gleason 3+3=6 disease, which may well seem concerning to many prostate surgeons. At a very early follow-up at 12 months, two men (25%) had biochemical recurrence and had residual prostate volumes of 6mL with positive biopsies. Both of the men with Gleason 3+4=7 disease and positive margins had biochemical recurrence at 12 months. The authors describe this as satisfactory, but these outcomes would not generally be considered acceptable in a comparable contemporary surgical series, particularly given the short follow-up.

We should applaud efforts to improve the functional outcomes of cancer surgery while preserving cancer recurrence rates, but unless a way can be found to accurately identify those men who truly do not have cancer near the fascia on the opposite side to the dominant lesion, such approaches may carry increased risks of local recurrence. Perhaps such procedures should be confined to only very low risk men, although the majority of such men do well without surgery.

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