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Introduction

This article aims to highlight various areas of endourology research that have had a direct effect on clinical practice in recent years.

Laser safety and technology

Perhaps inarguably, one of the biggest changes in endourology the last number of years has been the introduction of innovative laser technology into clinical practice. For this reason, we focus on laser technology and safety as the most important update in the management of urolithiasis in recent years.

Dr. Sero Andonian led a Canadian, prospective, randomized clinical trial comparing regular and Moses (Lumenis) modes of holmium laser lithotripsy.1 In their paper, Ibrahim et al investigated the performance of the Moses mode vs. regular mode of the Lumenis 120W laser generator (LP120H Moses) to assess stone retropropulsion, stone fragmentation/pulverization time, and procedural time during retrograde ureteroscopy. The Moses technology divides the holmium laser pulse into two adjacent peaks. The first peak separates the fluid ahead of the stone, creating a vapor bubble, hence the term Moses effect, which allows the second peak to be precisely delivered to the stone with less energy loss. Laser energy transmission is less dependent on fibre-to-stone distance with the Moses mode. In their study of 72 cases, Dr. Andonian and colleagues found that the Moses mode was associated with significantly shorter fragmentation/pulverization time (14.2 minutes vs. 21.1 minutes, p=0.03) and procedural time (41.1 minutes vs. 50.9 minutes, p=0.03) compared to regular (short-pulse) mode. There was less stone retropropulsion with the Moses mode (mean grade 0.5 vs. 1.0, p=0.01), which is an important consideration when treating ureteral stones. There was no difference between total energy used (10.8 kJ vs. 11.1 kJ, p>0.05), treatment success rates (88.4% vs. 83.3%, p>0.05), or complications (8.3% vs. 11.1%, p>0.05).

There have been several studies that assess the thulium fibre laser (TFL) use for stones and tissue uses. The TFL has changed our practices in that the technique of stone ablation has changed from less painting to more “pop-dusting.” Anecdotally, we feel that junior residents who are more novice ureteroscopists may be faster at learning laser lithotripsy, as they do not need to learn all the tips and tricks on how to perform fine movements with the ureteroscope to “paint” the face of the stone for fine dusting. Furthermore, we are expanding our clinical indications of ureteroscopy to treat larger or more complex stones less invasively rather than perform percutaneous nephrolithotomy. Other benefits are that the operating room is more pleasant, as the machine is quieter than the typical holmium laser, and the unit is very versatile, as it can be used with any standard electrical outlet. One caveat is to be aware of using high power in the ureter. Just as you would use no more than 10W cautiously with Holmium:yttrium-aluminum-garnet (Ho:YAG) in the ureter, one should not go above this level in the ureter using TFL. Similar wattage (power) from any laser equates to equal temperature rises and TFL is no different. Using higher power than 10W can result in higher temperatures that can lead to tissue ablation and stricture formation.2 Clinical use of the TFL has shown promising initial results and head-to-head trials will be likely be forthcoming in the next few years.3

Safety of the Ho:YAG laser has been assessed and a recent CUA best practice report (BPR) has been published by Bhojani et al.4 A study from 2015 using pig eyes showed that the Ho:YAG laser can produce eye injuries in the form of corneal abrasions.5 Researchers found that with a 272 micron laser fibre, the fibre needed to be within 5 cm of the pig’s eye in order to cause damage. No retinal or lens damage
was observed, only corneal damage. Regular prescription eyeglasses provided equivalent protection when compared to laser safety goggles. Accordingly, after reviewing this paper and other literature regarding the safety of Ho:YAG laser, the authors of the CUA BPR on Ho:YAG laser eye safety concluded that current evidence does not support mandatory safety eyewear for all operating room personnel and that standard eyeglasses are as protective as laser safety goggles.

### Analgesia: Minimizing opioid use

An Enhanced Recovery After Surgery (ERAS) protocol was developed by Gridley et al. for patients undergoing ureteroscopy with a goal to minimize perioperative opioid use and have an opioid-free protocol for postoperative pain management after discharge from hospital. The ERAS protocol is summarized in Table 1. The protocol included preoperative analgesia, intraoperative use of medications to help prevent bladder pain (belladonna and opium suppositories) and renal pain (ketorolac), a tiered approach to pain relief in the post-anesthesia care unit, and included no opioid prescription at discharge. Patients were discharged on acetaminophen, ibuprofen, an anticholinergic, and an alpha-blocker. Although this was not a randomized controlled trial, investigators found a significant difference pre-ERAS compared to the use of the ERAS protocol. Specifically, the mean total morphine milligram equivalent (MME) decreased from 60.1 MME to 7.7 MME (p<0.01) and the percentage of patients discharged with an opioid prescription decreased dramatically from 93% to 0% (p<0.01). There was no difference in calls for postoperative pain, nor were there differences in unplanned medical encounters due to uncontrolled pain. In this study, only 28.8% of patients in the ERAS group received opioids in the recovery room. There were no differences in quality-of-life outcomes between the two groups. Although this ERAS study has some limitations (for example, the use of tramadol), it is thought-provoking and could be used to develop similar opioid reduction strategies for endourological pain management.

### Alpha-blockers — Beyond medical expulsive therapy

The off-label use of preoperatively administered alpha-blockers to facilitate ureteroscopy for ureteral stones has been investigated. Pace and colleagues conducted a systematic review and meta-analysis of randomized controlled trials assessing alpha-blocker in this patient population. Alsaikhan et al pooled 11 randomized controlled trials comprising 1352 patients. They found there was a decreased need for ureteral dilation, with a 61% risk reduction (relative risk [RR] 0.39, 95% confidence interval [CI] 0.31–0.48, p<0.00001), increased ability to reach the stone(s) (RR 1.16, 95% CI 1.10–1.23, p<0.00001), improved stone-free rate (RR 1.17, 95% CI 1.08–1.26) at four-week followup, shorter operative time (weighted mean difference -6.05, 95% CI -10.17 to -1.93 min, p=0.004), and shorter hospital length of stay (mean difference [MD] -0.34, 95% CI -0.55 to -0.13 days, p=0.001).

The meta-analysis does leave some unanswered questions and the authors note that a large, adequately powered randomized controlled trial is recommended. Questions remaining include: What is the optimal duration of use? What is the effect of alpha-blockers prior to flexible ureteroscopy? Do alpha-blockers facilitate ureteroscopy in the pediatric population? The meta-analysis could also not control for size of the ureteroscope due to lack of data reported. Hopefully, these questions will be answered in future studies, although studying an off-label use of a medication does present some logistical challenges.

### Ureteric stents

As urologists, we often hear complaints from patients regarding symptoms related to ureteric stents. Although we sympathize with patients, many urologists may recommend ureteric stenting following ureteroscopy in the hopes of minimizing complications, such as strictures and return visits due to postoperative pain and hydronephrosis. A Cochrane review was recently published assessing the data to date regarding use of ureteral stents following uncomplicated ureteroscopy. This meta-analysis included 23 randomized controlled trials and 2275 patients, which represents the largest, most rigorously conducted analysis of this data to date. Unfortunately, the authors were unable to conduct pre-planned subgroup analysis of clinical data due to lack of more granular reporting in each study and there may have been selective reporting bias, as none of the 23 studies had been prospectively registered. The low-quality data

| Table 1. Analgesic components of the Enhanced Recovery After Surgery protocol for patients undergoing ureteroscopic stone surgery |
|---------------------------------------------------------------|
| **Preoperatively**                                           | Acetaminophen 650–1000 mg PO |
|                                                               | Gabapentin 100–300 mg PO     |
| **Intraoperatively**                                        | Belladonna/opium 30 mg PR   |
|                                                               | Ketorolac 300 mg IV          |
| **Post-anesthesia care unit**                               | First-line: Tramadol 50 mg PO as needed for pain rated 4–7/10 |
|                                                               | Second-line: Oxycodone 5 mg PO as needed for pain rated 4–7/10 |
|                                                               | Hydromorphone 0.25 mg IV for pain rated ≥8/10 |
| **Discharge**                                               | Acetaminophen 1000 mg every 8 hours for 7 days |
|                                                               | Ibuprofen 800 mg every 8 hours for 7 days |
|                                                               | Anticholinergic 10 mg XL daily prn for 10 days |
|                                                               | Alpha-blocker 0.4 mg daily for 10 days |
Conclusions

These were some of the more practice-changing papers we found in the literature over the last two years. It is great to see our field taking both major and minor steps to help us treat our patients more effectively.

Competing interests: Dr. Chew has been an advisory board member for ADVA-Tec, Ambu, Auris Surgical, Becton Dickinson, Boston Scientific, Cook Medical, Olympus, Storz Medical, Sanomotion, and The Ureteral Stent Company; has received grants/honoraria from ADVA-Tec, Ambu, Auris Surgical, Becton Dickinson, Boston Scientific, Cook Medical, Olympus, Storz Medical, Sanomotion, and The Ureteral Stent Company; hold investments in Auris Surgical, Sanomotion, and The Ureteral Stent Company; and has participated in clinical trials supported by ADVA-Tec, Boston Scientific, Cook Medical, Olympus, Storz Medical, and Sanomotion. Dr. Lantz Powers reports no competing personal or financial interests related to this work.

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Correspondence: Dr. Andrea G. Lantz Powers, Department of Urology, Dalhousie University, Halifax, NS, Canada; aglantz@dal.ca