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#### **CLINICAL INVESTIGATION**

**Normal Tissue** 

# HYPERBARIC OXYGEN TREATMENT OF CHRONIC REFRACTORY RADIATION PROCTITIS: A RANDOMIZED AND CONTROLLED DOUBLE-BLIND CROSSOVER TRIAL WITH LONG-TERM FOLLOW-UP

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Purpose: Cancer patients who undergo radiotherapy remain at life-long risk of radiation-induced injury to normal tissues. We conducted a randomized, controlled, double-blind crossover trial with long-term follow-up to evaluate the effectiveness of hyperbaric oxygen for refractory radiation proctitis.

Methods and Materials: Patients with refractory radiation proctitis were randomized to hyperbaric oxygen at 2.0 atmospheres absolute (Group 1) or air at 1.1 atmospheres absolute (Group 2). The sham patients were subsequently crossed to Group 1. All patients were re-evaluated by an investigator who was unaware of the treatment allocation at 3 and 6 months and Years 1–5. The primary outcome measures were the late effects normal tissue-subjective, objective, management, analytic (SOMA-LENT) score and standardized clinical assessment. The secondary outcome was the change in quality of life.

Results: Of 226 patients assessed, 150 were entered in the study and 120 were evaluable. After the initial allocation, the mean SOMA-LENT score improved in both groups. For Group 1, the mean was lower (p=0.0150) and the amount of improvement nearly twice as great (5.00 vs. 2.61, p=0.0019). Similarly, Group 1 had a greater portion of responders per clinical assessment than did Group 2 (88.9% vs. 62.5%, respectively; p=0.0009). Significance improved when the data were analyzed from an intention to treat perspective (p=0.0006). Group 1 had a better result in the quality of life bowel bother subscale. These differences were abolished after the crossover.

Conclusion: Hyperbaric oxygen therapy significantly improved the healing responses in patients with refractory radiation proctitis, generating an absolute risk reduction of 32% (number needed to treat of 3) between the groups after the initial allocation. Other medical management requirements were discontinued, and advanced interventions were largely avoided. Enhanced bowel-specific quality of life resulted. © 2008 Elsevier Inc.

Hyperbaric oxygenation, Controlled trial, SOMA-LENT, Late radiation injury, Quality of life.

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#### INTRODUCTION

Radiotherapy is a major nonoperative treatment and commonly used in the management of a number of different malignancies. During the past decade, significant developments in the delivery of radiotherapy have improved the efficacy and tolerance (1). Despite such advances, adverse effects continue to complicate its use (2, 3). These effects are commonly categorized as either acute effects, representing those that occur during or soon after radiotherapy completion, or late effects that manifest many months to several years later.

Acute toxicity is usually mild, frequently self-limiting, and often responds to brief interruptions in radiotherapy (3–5). Severe acute effects can lead to later excluded ones from "consequential" effects (6). Late toxicity is largely a function of the total radiation dose and fraction size and tends to be dose limiting in curative settings (7, 8). The resulting injuries are frequently refractory to a wide range of therapeutic interventions, can proceed to surgical removal of damaged organs, and are the cause of some mortality (2, 3, 9).

Late radiation proctitis is a particularly difficult condition to treat and for patients to live with (10–13). The reported incidence varies from 4% to 22% (5, 14), yet because of a frequent lack of recognition and insufficient long-term follow-up, its true incidence is unknown (14, 15). No recommended standard treatment exists, and current management is often unsatisfactory (11, 16). This shortcoming is readily apparent given the large number of medical and surgical therapies in common use (Table 1).

Hyperbaric oxygen (HBO) therapy has been used in the treatment of pelvic radiation injuries for several decades (Table 2) and has been reported to be beneficial (16–18). It

Table 1. Late radiation proctitis treatment options (in alphabetical order)

5-ASA Antidiarrheal agents Argon laser Cautery Corticosteroids Dilation and stenting Elemental diet Formalin Heat probe Hormonal therapy Hyperbaric oxygen therapy Iron supplementation Low-residue, low-fat diet Metronidazole Nd:YAG laser Pain control Pentosan Resection Replacement transfusion Short-chain fatty acids Sucralfate Surgical repair

Abbreviations: ASA = acetylsalicylic acid (aspirin); Nd:YAG = neodymium:yttrium-aluminium-garnet (laser) (Nd:Y<sub>3</sub>Al<sub>5</sub>O<sub>12</sub>).

has not, however, been studied in a sufficiently rigorous manner to determine its precise therapeutic effect. We conducted a multicenter, randomized, controlled, double-blind trial with crossover and long-term follow-up to evaluate the effect of HBO therapy for patients whose radiation proctitis had proven refractory to other interventions.

#### METHODS AND MATERIALS

#### Patients

Patients from the Instituto Nacional de Cancerologica, Mexico City, Mexico, the University of Pretoria Medical Centre, Pretoria, Republic of South Africa, Department of Underwater and Hyperbaric Medicine, Istanbul University, Istanbul Faculty of Medicine, Istanbul, Turkey, Wesley Medical Centre, Brisbane, Australia, and the Royal Hobart Hospital, Tasmania, Australia were enrolled in the trial. Each participating center's institutional review board approved the study protocol. Referring physicians agreed to participate as blinded assessors. The trial registration numbers were NCT00134628 and ISRCTN85456814.

Patients were eligible for enrollment if they had undergone pelvic radiotherapy and had subsequently developed evidence of rectal late radiation tissue injury. The diagnosis had to have been present for ≥3 months and to not have responded sufficiently to other therapies. Eligibility screening confirmed the absence of unacceptable patient-specific risks to HBO therapy. All patients or their surrogate provided written informed consent before enrollment. On patient enrollment, the best supportive care was maintained.

Before beginning treatment, patients were evaluated with the late effects normal tissue-subjective, objective, management, analytic (SOMA-LENT) scale, an anatomic-specific morbidity scoring system (19). It provides an ascending order of severity of radiation-induced complications. It is particularly well suited to multicenter trials, because of its standardized application, reproducibility, and accuracy. A standardized clinical assessment was also included with both screening tools conducted by a physician unaware of the allocation. Patients also completed the Expanded Prostate Cancer Index Composite (20) quality of life (QOL) instrument at this time and at every other follow-up stage.

#### Randomization

Biostatisticians at the University of South Carolina generated the randomization sequence, which was uploaded into, and concealed within, the study database software. The patients were randomly assigned (1:1) to receive HBO or normobaric air, using a "blocking" process. The block size was four and was equally stratified with two of each treatment options (A or B). The randomization sequence became available to the unblinded local principal investigator only on irretrievable entry of each patient's demographic information, medical history, and clinical characteristics. Group 1 (active treatment) was randomized to receive 2.0 atmospheres absolute (ATA) oxygen. Group 2 (sham) patients were randomized to receive 1.1 ATA air.

#### Treatment procedure

Group 1 was treated with 100% oxygen at 2.0 ATA for 90 min, once daily, five times weekly. Group 2 were treated with 21% oxygen (normal air) at 1.1 ATA, once daily, five times weekly. For patient blinding purposes, Group 2 patients underwent a brief compression to 1.34 ATA at the beginning of each treatment. The chamber was then slowly decompressed from 1.34 to 1.1 ATA.

Table 2. Reported hyperbaric oxygen dosing and outcomes for radiation proctitis

		Н	yperbaric treatment		
Investigator	Patients (n)	Pressure (ATA)	Time (min)	Treatment sessions (n)*	Overall improvement (%)
Bouachour et al. (31), 1990	8	2.5	90	$80 \pm 10$	75
Charneau et al. (28), 1991	1	2.5	?	80	Healed
Nakada et al. (35), 1993	1	2.0	90	30	Healed
Hamour et al. (36), 1996	1	2.5	90	49	Healed
Feldmeier et al. (37), 1996	7	2.4	90	3-50 (24)	57
Woo et al. (38), 1997	18	2.0	90	12–40	>50
Warren et al. (39), 1997	14	2.0-2.5	90-120	?	59
Ugheoke et al. (40), 1998	8	2.5	90	20-40 (28)	62.5
Carl et al. (41), 1998	2	2.4	90	38–40 (39)	50
Gouello et al. (42), 1999	36	2.5	90	Mean 67	56-65
Kitta et al. (43), 2000	4	2.0	60	30-60 (38)	75
Bem et al. (44), 2000	2	2.4	90	60	100
Roque et al. (45), 2001	6	2.5	90	20-60 (37)	85
Mayer et al. (46), 2001	7	2.2-2.4	60	20–60 (33)	85
Boyle et al. (47), 2002	19	2.0	120	27–80 (59)	68
Jones et al. (48), 2006	10	2.0-2.5	90	36–41 (40)	>70
Dall'Era et al. (49), 2006	27	2.4	90	29–60 (36)	48
Fink et al. (50), 2006	4	2.4	90	20–50 (33)	50
Girnius et al. (51), 2006	9	2.5	90	22-80 (58)	78
Nakabayashi et al. (52), 2006	1	2.4	90	40	Healed
Marshall <i>et al.</i> (53), 2007	65	2.36	90	30-60	25-73

Abbreviation: ATA = atmospheres absolute.

Group 2 patients remained for the sum of the time taken to treat the Group 1 patients. Reassessment, after 30 treatment sessions, was undertaken by the referring physician, who remained unaware of the allocation. Ten additional treatment sessions were provided to selected patients, depending on the individualized responses. Patients repeated their QOL survey and were screened to determine the effectiveness of the blinding process. Unblinding took place at this point.

Those who had been allocated to Group 1 were entered into follow-up, with repeat evaluations scheduled at intervals of 3 and 6 months and Years 1–5. For Group 2, all but 3 accepted crossover to the active treatment arm.

#### Data collection at inclusion

Once a patient was enrolled, their local principal investigator collected the following data: age and gender; comprehensive medical history; current medications and any history of tobacco use; cancer-related history, including tumor type, location, stage, and treatment; and late radiation proctitis signs and symptoms, including treatment sessions to date.

#### Statistical analysis

The primary outcome was a change in the SOMA-LENT (Fig. 1) score, a numeric variable measured at all periods. Four other numeric values were derived from a QOL survey completed by patients in conjunction with their clinical evaluations. From this survey, using the Expanded Prostate Cancer Index Composite Bowel Domain, the Bowel Function and Bowel Bother subscales were obtained. Also obtained were the physical and mental results using the SF-12 General Health Function Survey. The SOMA-LENT score was analyzed using a repeated measures model containing patient type, period, their interaction, and six covariates: gender, tobacco use, external beam radiotherapy and brachytherapy, interval

between radiotherapy and symptoms, interval between symptoms and treatment, and country of residence.

A sixth, ordinal categorical outcome, was the clinical evaluation measured at all periods, except at initialization. The evaluations made immediately after completion of the initial treatment allocation and crossover were coded as healed, significant improvement, modest improvement, or no improvement. For the remaining periods, they were coded as healed, improved, unchanged, or recurrence. For analysis purposes, these evaluations were dichotomized. After the initial treatment allocation and crossover, healed, significant improvement, and modest improvement were collapsed into one category and no improvement and recurrence into the other. For the follow-up evaluations, healed and improved were collapsed into one category and no improvement and recurrence into the other. The outcomes were compared for the two patient types using Fisher's exact test and logistic regression analysis containing the same variables as the repeated measures model for SOMA-LENT. Additionally, a Jonckheere-Terpstra test for trend was used with the original calculations.

#### **RESULTS**

A total of 226 patients were assessed for eligibility. Of these 226 patients, 76 were excluded and 150 enrolled. Of the 150 patients, 120 completed the protocol (Fig. 2). At 1 year, 5 patients (4%) had died and 9 (8%) had been lost to follow-up.

#### Descriptive statistics

Data were available for 120 patients. The minimal followup period for all patients was 1 year (average, 2.09). Of the 120 patients, 106 (88.33%) were women, and 101 (84.17%)

<sup>\*</sup> Average number of treatment sessions in parentheses.

## **SOMA LENT scoring system for radiation proctitis**

TITIS		NT NAME:			HORTIS IV
HORTIS I.E	)SIGNA	TURE:		DATE: _	
GRADE 1	GRADE 2	GRADE 3	GRADE 4	<u> </u>	-
				SCORE	FACILITY
Occasional urgency	Intermittent urgency	Persistent urgency	Refractory		CODE:
Optasional	Intermittent	Persistent	Refractory		Scoring Instructions:
Occasional	Intermittent	Persistent	Refractory		Score the 14
2 - 4 per day	4 - 8 per day	> 8 per day	Uncontrolled diamhea		SOM paramet with 1-4 and
Occasional & minimal	Intermittent & tolerable	Persistent & intense	Refractory & excruciating		total all 14 to
					generate the
Occult	Occasionally >2/week	Persistent/daily	Gross hemonhage	ļ	1st LENT Score
Superficial ≤ 1 cm²	Superficial > 1 cm <sup>2</sup>	Deep ulcer	Perforation, Fistulae		(Score ≈ 0 if
> 2/3 normal diameter with dilation	1/3 - 2/3 normal diameter with dilation	< 1/3 normal diameter	Complete obstruction		there are no toxicities)
				7	
Op:asional, <u>≤</u> 2 antidiarrheals/week	Regular, > 2 antidiamheals/week	Multiple. > 2 antidiamheals/day	Surgical intervention/ Permanent colostomy		1st LENT Score
Occasional non-narcetic	Regular non-narcotic	Regular narcotic	Surgical intervention		Divide the 1st LENT Score
Stool softener, iron therapy	Occasional transfusion	Frequent transfusions	Surgical intervention / Permanent colostomy		by 14 to provide the 2nd LENT
Diet modification, stool softener	Occasional steroids	Steroids per enema, hyperbaric oxygen	Surgical intervention / Permanent colostomy		Score
Diet modification	Occasional dilatation	Regular dilatation	Surgical intervention		2nd LENT Score
Oppasional use of incontinence pads	Intermittent use of incontinence pads	Persistent use of incontinence pads	Surgical intervention / Permanent colostomy		
				Y/N Dat	
3					
		non			
		ine			
	HORTIS I.E  GRADE 1  Occasional urgency Occasional Occasional 2 - 4 per day Occasional & minimal Occult Superficial ≤ 1 cm² > 2/3 normal diameter with dilation  Occasional = 2 antidiameter with dilation  Occasional non-narcotic Stool softener, iron therapy Diet modification, stool softener Diet modification Occasional use of incontinence pads  Assessment of lumen and Assessment of wall thickn Assessment of wall thickn Assessment rectal compli	HORTIS I.D. SIGNA  GRADE 1 GRADE 2  Occasional urgency Intermittent urgency Occasional Intermittent  2 - 4 per day 4 - 8 per day Occasional Intermittent & tolerable  Occusional & minimal Intermittent & tolerable  Occusional & Tom² Superficial > 1 cm²  > 2/3 normal diameter with dilation  Occasional, ≤ 2 antidiamheats/week  Occasional non-narcotic  Stool softener, iron therapy  Diet modification, stool softener  Diet modification, stool softener  Diet modification Occasional transfusion  Occasional use of incontinence pads  Assessment of lumen and peristalsis Assessment of wall thickness, sinus and fistula formal Assessment rectal compliance	SIGNATURE:  GRADE 1 GRADE 2 GRADE 3  Occasional urgency Intermittent urgency Persistent urgency Occasional Intermittent Persistent Occasional Intermittent Persistent 2 - 4 per day 4 - 8 per day > 8 per day Occasional Intermittent Persistent  2 - 4 per day 4 - 8 per day > 8 per day Occasional Intermittent & tolerable Persistent & intense  Occult Occasionally >2/week Persistent & intense  Occult Occasionally >2/week Persistent/daily Superficial ≤ 1 cm² Superficial > 1 cm² Deep ulder  > 2/3 normal diameter with dilation  Occasional, ≤ 2 Regular, > 2 antidiamhea/s/week antidiamhea/s/week Occasional non-narcotic Regular non-narcotic Regular narcotic  Stool softener, iron therapy Diet modification, stool softener Diet modification Occasional steroids Steroids per enema, hyperbaric oxygen Diet modification Occasional dilatation Regular dilatation Occasional use of Intermittent use of Persistent use of incontinence pads  Assessment of lumen and peristalsis Assessment of lumen and peristalsis Assessment of wall thickness, sinus and fistura formation Assessment of wall thickness, sinus and fistura formation Assessment of wall thickness, sinus and fistura formation	Cocasional urgency	SIGNATURE:   GRADE 1   GRADE 2   GRADE 3   GRADE 4

Fig. 1. Late effects normal tissue-subjective, objective, management, analytic (SOMA-LENT) scoring system for radiation proctitis.

reported never having smoked. Because of the small number of current (n = 8) and former (n = 11) smokers, the tobacco variable was dichotomized into ever/never. Of the 120 patients, 11 (9.17%) were from Australia, 85 (70.83%) from Mexico, and 12 (10.00%) from both South Africa and Turkey. The baseline comparisons of the covariates for the two groups resulted in no significant differences, indicating that the randomization process had worked well. The patient demographics and clinical characteristics are detailed in Table 3 (appears online only at www.redjournal.org). The mean SOMA-LENT values for the two patient types at each period are displayed in Fig. 3. The mean SOMA-LENT score decreased considerably between the initial value and completion of HBO therapy in Group 1, with a much smaller change in Group 2. For the latter group, however, a substantial decrease occurred after crossover, when they received HBO therapy.

### Numeric outcomes

*SOMA-LENT score*. Adjusting for covariates, a significant (p < 0.0001) decrease (improvement) occurred in Group 1 of

5.00 (95% confidence interval, 3.96–6.03), as well as a significant (p < 0.0001) decrease in Group 2 of 2.61 (95% confidence interval, 1.51–3.70) after completion of the initial allocation. The decrease was greater in Group 1 than in Group 2 (p = 0.0019). At initialization, no difference was detected between the two groups (p = 0.5597). However, after the initial allocation, Group 1 had significantly (p = 0.0150) lower average scores than Group 2, with an estimated difference of 1.93 (95% confidence interval, 0.38–3.48). After completion of the crossover, no differences were detected (p = 0.6594). The mean scores remained relatively stable through 1 year and showed a trend to additional and sustained improvement through Year 5.

Clinical evaluation. The frequencies for clinical evaluations are given in Table 4. The most notable result was after completion of the initial allocation, at which 56 (88.9%) of the 63 patients in Group 1 were assessed to have either healed or had some improvement, and 35 (62.5%) of the 56 patients in Group 2 were assessed to have had at least some improvement. Fisher's exact test (p = 0.0009) and logistic regression

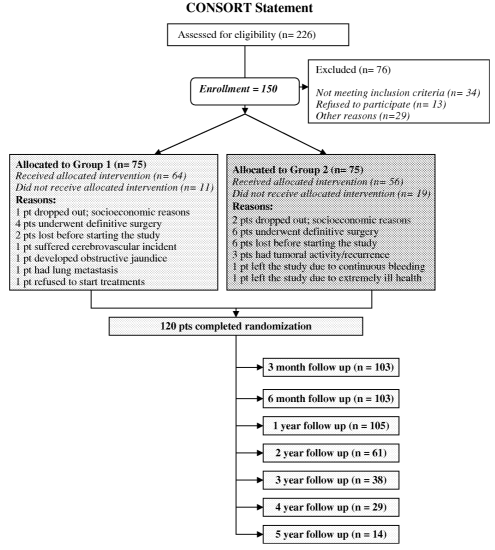


Fig. 2. Consolidated Standards of Reporting Trials (CONSORT) statement.

analysis (p = 0.0011) both indicated that Group 1 had a significantly greater proportion of healing/improvement at that time. For logistic regression analysis, the corresponding odds ratio was 5.93 (95% confidence interval 2.04–17.24). From this, we estimated that Group 1 was about six times more likely to have an evaluation that indicated at least some type of improvement than was Group 2. Furthermore, the Jonckheere-Terpstra test for trend was significant (p = 0.0008), indicating that better outcomes were more common in Group 1. On the basis of the clinical evaluation outcomes, an absolute risk reduction of 0.32 (32%) was generated, resulting in a number needed to treat of 3.

From an intention to treat perspective, we considered what would have happened if (1) all those for whom we had no results had had improvement, (2) all those for whom we had no results had not had improvement, and (3) for each patient type, one-half of those for whom we had no results had improvement and one-half had not. In all cases, the results still indicated that Group 1 had a significantly greater

proportion of improvement than did Group 2 (p = 0.0057, p = 0.0007, and p = 0.0036, respectively).

Quality of life. Marked improvement was noted in the bowel-specific QOL assessment for Group 1 after treatment but not for Group 2 (14% for Bowel Bother and 9% for Bowel Function vs. 5% and 6%, respectively). After crossover, Group 2 showed notable improvement, with an increase to 13.6 for bowel bother and 10% for bowel function. Both groups showed additional improvement at 1 year. For the bowel bother subscale, a significant improvement was seen between initialization and randomization in Group 1 (estimated change, 14.14; p = 0.0007, adjusting for covariates), but not in Group 2 (estimated change, 5.75; p = 0.1521). However, Group 2 experienced a significant improvement after crossover (estimated change, 14.27; p = 0.0002). The scores for both groups were stable or tended to improve further throughout follow-up. Similar trends were seen in the bowel function subscale. No differences were observed in the general well-being assessment.

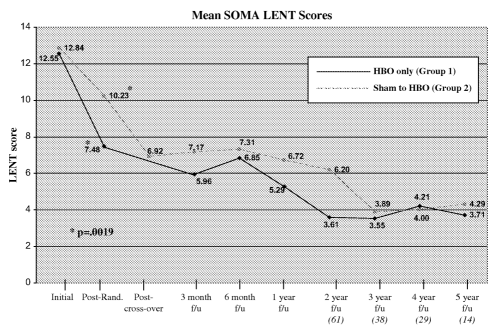


Fig. 3. Mean late effects normal tissue-subjective, objective, management, analytic (SOMA-LENT) scores. HBO = hyperbaric oxygen.

#### Patient beliefs

Of the 120 patients, 72 (33 in Group 1 and 39 in Group 2) were surveyed to determine which randomization allocation they had received. In Group 1, 20 said "HBO," 1 said "sham," and 12 "could not say." In Group 2, these numbers were 23, 2, and 14. A chi-square test detected no relationship (p = 0.9058) between the patient opinions and what they had actually received. When patients who "could not say" were ignored, a Kappa statistic was p = 0.0299, indicating essentially no agreement beyond chance.

#### Harms

Consistent with hyperbaric practice, ear pain/ear discomfort (ear barotrauma) was the most common complaint. Ear barotrauma represents the clinical manifestation of an imbalance of pressure between the external and middle ear spaces. It is usually limited to the tympanic membrane, occasionally involves the middle ear, and only rarely involves the inner ear. Nineteen patients (15.8%) complained of ear pain or discomfort. The otologic examination was unremarkable in 11, 7 had tympanic membrane changes consistent with barotrauma, and 1 had both tympanic membrane injury and middle ear effusion. Decongestants were effective in 8 patients, 7 underwent ventilation tube placement, and 4 did not require treatment. One patient (0.8%) complained of sinus barotrauma and was successfully treated with decongestants.

Four patients (3.3%) experienced transient myopia. This is a poorly understood process and although thought to represent an oxidative stress-induced temporary alteration in the shape of the lens (21), its exact mechanism remains obscure.

Two patients (1.7%) complained of confinement anxiety. One was treated with reassurance alone; the other required mild sedation. No cases of acute central nervous system oxygen toxicity occurred. None of these harms compromised

a patient's participation in the study, and all patients completed their prescribed treatment course.

### DISCUSSION

Radiation proctitis is a common unfortunate complication of pelvic radiotherapy (22). Its reported incidence ranges from 4% to 22% (5, 7, 14) and can reach 36% after combination external beam radiotherapy and brachytherapy (23). More severe forms, some of which are life-threatening, have been reported to range from 4.3% to 22% (14, 24) with resulting mortality rates of 2–8% (3, 7, 24).

Most late cases occur within 3 years of radiotherapy completion, although latencies in excess of 10 years are not uncommon (14, 22). The natural history of late radiation proctitis is unpredictable. Minor symptoms can resolve either spontaneously (4) or with conservative management (2, 25). Other seemingly minor symptoms will prove refractory to standard care, resulting in disease progression despite increasingly aggressive interventions (24), and new forms of this complication can evolve (22). Minor complaints of pain and bleeding, therefore, cannot be characterized as harmless manifestations. Serious manifestations can necessitate high-risk surgery; high risk because tissues within the operative site might have been rendered hypoxic and poorly able to support oxygen-dependent wound repair. Ultimately, and having survived cancer, some patients will die of these complications (3, 7, 24).

The clinical presentation can involve any combination of tenesmus, urgency, diarrhea, constipation, sphincter dysfunction, mucoid or bloody discharge per rectum, frank bleeding, and ulceration, which can be localized, diffuse, or full thickness. The mucosa can appear granular, friable, edematous, and pale, with prominent submucosal telangiectatic

Table 4. Frequencies of clinical evaluations by patient type

Evaluation point	Clinical evaluation findings	Group 1	Group 2
Randomization*	Healed	5	0
	Significant improvement	24	15
	Moderate improvement	27	20
	No improvement	7	21
Crossover	Healed	1	3
	Significant improvement	0	33
	Moderate improvement	1	11
	No improvement	1	6
3-mo	Healed	5	2
	Improved	31	26
	Unchanged	18	18
	Cancer recurrence	1	2
6-mo	Healed	4	3
	Improved	30	24
	Unchanged	19	17
	Cancer recurrence	2	4
1-y	Healed	5	2
•	Improved	32	30
	Unchanged	17	16
	Cancer recurrence	1	2
2-y	Healed	6	1
•	Improved	21	12
	Unchanged	8	11
	Cancer recurrence	1	1
3-у	Healed	2	3
•	Improved	15	12
	Unchanged	3	3
	Cancer recurrence	0	0
4-y	Healed	2	2
•	Improved	12	10
	Unchanged	0	3
	Cancer recurrence	0	0
5-y	Healed	1	0
•	Improved	4	6
	Unchanged	1	0
	Cancer recurrence	0	1

 $<sup>^{*}</sup>$  p Values comparing groups after randomization were 0.0009 for Fisher's exact test, 0.0011 for logistic regression analysis, and 0.0008 for Jonckheere-Terpstra test for trend.

vasculature. Pain is common, ranging from occasional and minimal to refractory and excruciating.

The histologic findings can include microvascular compromise, endothelial cell degeneration, and formation of fibrin plugs (26). Submucosal fibrosis and obliteration of small blood vessels is additional evidence of late radiation injury. This process is usually progressive and irreversible. Computed tomography can demonstrate wall thickening, edema, ulcers, stricture, and fistula (27).

The medical treatment is not well defined and, in the absence of recommendations, management is often unsatisfactory (3, 8, 12, 22). One should do everything possible to avoid disease progression, however, because abdominopelvic operations (unavailable in the presence of perforation, obstruction, and fistula) within or through irradiated tissues are fraught with complications (8, 28).

High failure rates with conventional treatment led to the use of HBO therapy. Its beneficial effect, involving mandibular osteoradionecrosis, was first reported in 1973 (29). Resulting pathologic evidence of a progressive and

obliterative endarteritis in mandibular osteoradionecrosis contrasted sharply with earlier assumptions of an osteomyelitic-like process (30). The finding that HBO therapy induced angiogenesis, suggested a disease-modifying mechanism, in contrast to more conventional medical and surgical therapies directed at relief of symptoms (16, 17).

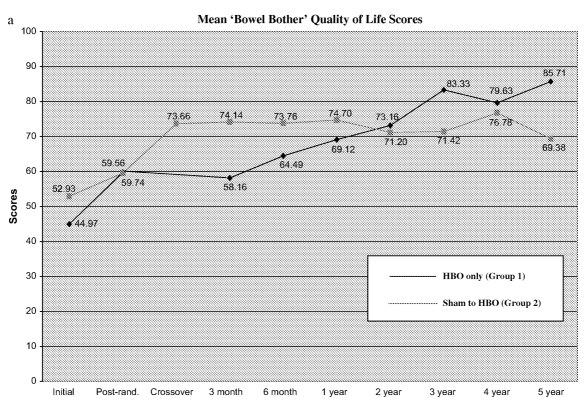
Hyperbaric oxygen therapy was first reported to have efficacy in the treatment of late radiation proctitis in 1990 (31). Since then, numerous studies have been published (Table 2). In most instances, they represented small series or single case reports, did not use a specified toxicity scale, and lacked sufficient follow-up. However, the results from this accumulated work do suggest that HBO therapy is likely to be beneficial (16, 18).

We used SOMA-LENT scoring as a primary outcome measure. This numeric evaluation of radiation morbidity is simple, widely applicable, reproducible, and provides an ascending order of severity (19). Given that several different physicians would evaluate outcomes in this multicenter study, such uniform scoring was considered essential. The radiation proctitis SOMA-LENT process scores symptoms on a severity scale of 1–4 for each of five possible symptoms and three related objective clinical signs. Six management options, scored in increasing complexity, represent the final scoring element. The analytic measures used during the diagnostic workup can be recorded but are not scored.

Often, the outcome assessment is a function of clinical impression alone. This, however, opens evaluations to differences in interpretation and has the potential for bias. We elected to include this approach as a second primary outcome measure. Perhaps not surprisingly, the resulting percentage of clinical assessments determined as healed was lower than those reported in several previous studies. The specificity of the SOMA-LENT scale is such that an excellent healing response does not always result in a score of 0 (healed). A final response score of 2-3 might reflect a patient who, on presentation had a score of 15 for ulceration, intense pain, and persistent bleeding, required treatment with narcotics, occasional transfusions, and steroids, and whose post-treatment status became one of diet modification, twice-daily stool frequency, and an occasional non-narcotic analgesic. The clinical impression of this case would be one of "healed" by many. In the present trial, however, the clinical assessor also conducted each SOMA-LENT analysis. Recognizing that the score was not 0, the assessor might have been inclined to categorize the clinical outcome as something less than healed (e.g., significantly improved).

The effect of HBO therapy, scored through the SOMA-LENT process, throughout the 5-year study period is shown in Fig. 3. Although the number of patients at Years 2–5 was 58%, 36%, 27%, and 13% of those at Year 1, respectively, a clear trend was seen toward continued and enduring healing.

A patient's perception of how effective a particular treatment is now represents one important element of the modern application of evidence-based medicine (32). The QOL effect of eliminating pain, minimizing hemorrhage, and normalizing stool frequency is obviously important. This effect was



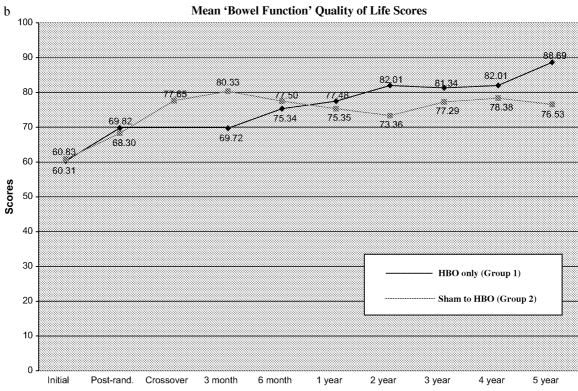


Fig. 4. (a) Bowel bother and (b) bowel function quality of life scores. HBO = hyperbaric oxygen.

evidenced by a significant improvement in the QOL recorded after receipt of HBO therapy in each group. The values continued to improve in Group 1 throughout the 5-year study period for bowel bother and bowel function. In Group 2, bowel bother continued to improve, and bowl function stabi-

lized at its 1 year value throughout the remainder of the study (Fig. 4).

One final observation of some importance was an association between failure to respond and a finding of local recurrence or residual tumor. Three patients were diagnosed

with recurrence during the treatment phase. Eleven others were diagnosed during follow-up, for a recurrence rate of 11.7%. The SOMA-LENT scores in these patients had either remained elevated or improved, only to acutely deteriorate, by an average of 9 points (range, 4–17), by the time the recurrence was diagnosed.

In our study, approximately 45% of those patients without a treatment response were diagnosed with local recurrence. This finding argues for a measured approach to hyperbaric dosing. Ordering an initial hyperbaric course of more than 40 sessions is inadvisable. If little or no subsequent improvement occurs, workup for cancer recurrence should occur before any further hyperbaric treatments.

Hyperbaric oxygen therapy was well tolerated and its safety profile proved encouraging. These findings are consistent with standard practice, with hyperbaric medicine considered low risk. Predictably, no cases of oxygen toxicity developed. This was one of our study's safety goals, with the resulting treatment pressure selection of 2.0 ATA.

A patient's perception of how well, or otherwise, a specific therapy effects their daily living and overall QOL has only recently been recognized as an important outcome measure (32). In our study, patients considered HBO therapy to have an important positive effect on their QOL when measured against their primary complaint.

When numerous therapeutic options exist for a given condition, responsible resource expenditure assumes increasing importance. Although hyperbaric medicine's costs are not insignificant, its employment has resulted in an overall lowering of a patient's total healthcare financial burden (33, 34). Much of this cost reduction is achieved by avoiding repeated hospi-

talizations and surgeries, because greater disease resolution rates are effected. Such savings support a preference for disease-modifying interventions rather than those directed at relief of symptoms. The immediate and enduring effect of HBO therapy on the resolution or reduction in the degree of radiation proctitis would be expected to have a corresponding positive effect on the overall cost of care. Although we did not incorporate an economic analysis in this trial, several assumptions can be made. First, because disease progression is not uncommon (2, 9), avoiding it would be expected to result in a corresponding decrease in the healthcare costs necessary to manage advancing degrees of morbidity and the costs associated with management failure. Second, a reduction in disease severity, or its resolution, likewise would reduce the subsequent costs. Using the example of the mean improvement in SOMA-LENT change at 1 year in our trial, an index patient's requirements would change from repeated rectal examinations, regularly administered narcotics, multiple daily antidiarrheal agents and steroid enemas to occasional antidiarrheal agents, diet modification, and perhaps a stool softener. The financial implications related to this change in medical management are readily calculable.

#### **CONCLUSION**

The results of our study have shown that the provision of HBO therapy for patients with chronic refractory radiation proctitis resulted in significantly improved and enduring healing responses and enhanced QOL. Our results support the role of HBO therapy for soft-tissue radionecrosis.

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#### **APPENDIX**

The clinical evaluation team included Dr. Fulya Yaman Agaoglu, Dr. Ridvan Avul, Dr. Robyn Cheuk, Dr. Yavuz Dizdar, Dr. Aaron Gonzalez, Dr. Susan Houldsworth, Dr. Esra Kaytan, Dr. Everine Klopper, Mr. Robert Lincacre, Dr. James Mackean, Dr. Robbie de Meulenare, Dr. Aida Mota, Dr. Gonzalo Montalvo, Dr. Browwyn Mueller,

Dr. Binnur Pinarbasi, Dr. Ken Purdie, Dr. Marlene Soares, Dr. John Stephenson, Dr. Colin Tang, Dr. Cherian Varughese, Dr. Albert Verbeek, Dr. Margaret Wallington, and Dr. Alida Wolvaardt. The study monitors included Ms. Guillermina Silva, Lic.Enf., Dr. Baris Pekicten, Mr. Stephen Goble, and Ms. Surita Fitchat, R.N.

Table 3. Patient demographics

	Diabetes mellitus Hypertension Transfusions	No	<sup>o</sup> N	°N	°Z	°Z	°Z	Yes	Yes	°Z
	Hypertensi	No	No	No	Š	No	No	No	No	No
	Diabetes mellitus	No	N <sub>o</sub>	No	o O	N <sub>o</sub>	o O	No	N <sub>O</sub>	o N
	Tobacco	Never	Never	Never	Never	Never	Never	Never	Never	Never
	$\begin{array}{c} \text{Previous} \\ \text{LENT} \\ \text{treatment}^{\dagger} \end{array}$	2,5	2	2	2	2	2	2, 12 (Diet)	1, 2, 12 (Sucralfate)	61
	s LENT presentation	Diarrhea, hemorrhage, stricture	Hemorrhage	Diarrhea, hemorrhage	Diarrhea, hemorrhage	Pain, hemorrhage	Hemorrhage	Pain, hemorrhage, ulceration, stricture	Hemorrhage	Hemorrhage
Time to	LENT diagnosis $(mo)^*$	12	16	12	16	13	13	10.5	41	9
	RT/dosage	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachtherapy, 3,031.75 cGy	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	2,007 COy X-ray, 4,600 cGy (200 cGy x 23 fractions) Brachytherapy, 3,344 cGy	2,574 COy X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy, 3,961 cGy	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy, 3,605 cGy cGy cGy cGy x 25	X-ray, 5,000 cGy (200 cGy x 16 + 300 cGy x 8) Brachytherapy, 2 571 cGy	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy, 2,000 cGy 2,000 cGy x 25	2.04-0 COy X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy, 3.841 cGy
Cancer treatments	Chemotherapy (type/dose)	No	Š	N	N O	N	Š	S N	N <sub>o</sub>	Cisplatin/ 250 mg
	Surgery (type)	No	Hysterectomy + BSO	No	Hysterectomy + BSO	N	N	Extrafascial hysterectomy	N	° Z
	Cancer type/stage	SCC/IIIb	AC/IIb	SCC/IIb	AC/Ib2	SCC/IIIb	SCC/IIa	AC/IIb	SCC/Ib2	SCC/IIIb
	Tumor location	Uterine cervix	2/26/1999 Uterine cervix	12/8/1999 Uterine cervix	11/26/1998 Uterine cervix	Uterine cervix	11/4/1999 Uterine cervix	2/11/2000 Uterine cervix	8/11/1995 Uterine cervix	8/24/1999 Uterine cervix
,	Cancer diagnosis date	3/16/1998	2/26/1999	12/8/1999	11/26/1998	6/9/1999	11/4/1999	2/11/2000	8/11/1995	8/24/1999
	Gender	щ	II.	<u>r</u>	II.	Ľ.	II.	ш	IT.	IL,
	Patient ID (	PROC 001B	PROC 002B	PROC 003A	PROC 004A	PROC 005A	PROC 006A	PROC 007B	PROC 008B	PROC 009A

Hemorrhage 2 Never No No No	nage 2 Never No No No		nage, wall I	rage, wall 1 ges Mic) Mic) Aic)	1 4 Never No Yes 2, 12 (Diet, Never Yes No steroid enema)	1 4 Never No Yes 2, 12 (Diet, Never Yes No steroid enema) 2 Never No No	wall 1  4 Never No Yes litis steroid enema)  1, 2 Never No No 11, 12 (Sucralfate Never No No stitis and cystifis after RT with dimethyl sulfoxide)	wall 1  4 Never No Yes  1, 2 12 (Diet, Never Yes No Sets of Strings and cystitis after RT with dimethyl Sulfoxide)  11, 12 (Sucralfate Never No No Sets of Steroids, Bicap)  2 Never No No Sets of Steroids, Bicap)	wall 1  4 Never No Yes  11, 12 (Sucralfate Never No No  11, 12 (Sucralfate Never No No  12, 11, 12  13, 2, 11, 12  14, 12, Never No No  15, 2, 12, Obiet, Never No No  16, 3, 11, 12  17, 12 (Sucralfate Never No No  18, 2, 11, 12  18, 11, 12  19, 11, 12  10, 11, 12  11, 13  11, 14  11, 15  11, 1
	13 Hemorrhage 2	14 Hemorrhage, wall 1 changes	(fibrotic)		(fibrotic)  Hemorrhage Pain, hemorrhage, Unspecific chronic colitis	Hemorrhage Pain, hemorrhage, Unspecific chronic colitis Diarrhea, pain, hemorrhage, stricture, mild chronic colitis	Hemorrhage Pain, hemorrhage, Unspecific chronic colitiis biarrhea, pain, hemorrhage, stricture, mild chronic colitiis 5 Hemorrhage,	Hemorrhage Pain, hemorrhage, Unspecific chronic colitis biarrhea, pain, hemorrhage, stricture, mild chronic colitis Chronic cystitis Chronic cystitis Diarrhea, pain, hemorrhage, ulceration	Hemorrhage Pain, hemorrhage, Unspecific chronic colitis biarrhea, pain, hemorrhage, stricture, mild chronic colitis Chronic cystitis Chronic cystitis Uleration biarrhea, pain, hemorrhage, ulceration chronic mild chronic mild chronic opitis
fractions) Brachytherapy, 3,600 cGv	13	00 cGy 14 c 25 apy,		0 cGy 14 (16 apy,	4	3y . 35 . 14	3y 35 Pa He Dii	3y 35 Pa He He Jiy 35 Da Jiy 35 Pa Jiy 35 Pa Jiy 19 Di J	y 14 He y 35 Pa y 14.5 He y 14.5 He y 11.5 Pa
		Cisplatin/ 420 mg	Š					ini/ mg	
	°Z	°N	No		Hysterectomy + BSO	Hysterectomy + BSO BSO Radical hysterectomy and pelvic kymphadenectomy	Hysterectomy + BSO BSO Radical hysterectomy and pehvic fymphadenectomy Abdominal hysterectomy	Hysterectomy + BSO Radical hysterectomy and pelvic hymphadenectomy Abdominal hysterectomy	Hysterectomy + BSO BSO Radical hysterectomy and pelvic hymphadenectomy No No No
	SCC/IIIb	SCC/IIa	SCC/IIb		SCC/Ib1	SCC/Ib1	SCC/Ib1 SCC/Ib1	SCC/Ib1 SCC/IIb SCC/IIIb	SCC/Ib1 SCC/IIb SCC/IIIb SCC/IIIb
cervix	10/24/1994 Uterine cervix	12/1/1999 Uterine cervix	1/19/2000 Uterine cervix		4/21/1998 Uterine cervix	4/21/1998 Uterine cervix 4/13/2000 Uterine cervix	4/21/1998 Uterine cervix 4/13/2000 Uterine cervix 8/20/1998 Uterine cervix	4/21/1998 Uterine cervix 4/13/2000 Uterine cervix 8/20/1998 Uterine cervix 11/8/1999 Uterine cervix	4/21/1998 Uterine cervix 4/13/2000 Uterine cervix 8/20/1998 Uterine cervix cervix 6/29/2000 Uterine cervix
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Table 3. Patient demographics (continued)

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(Continued)

mellitus Hypertension Transfusions Yes å å ž å å ž ν̈́ Yes Š Yes ŝ ž Yes ν̈́ Š ν̈́ Tobacco Diabetes Yes Yes Yes Yes å ž ž ŝ ν̈́ Current use Current use Past use Never Never. Never Never. Never Never. nse 12 (Pentoxyphiline, tocopherol) 2, 12 (Diet) Previous LENT reatment 2, 11 2, 5 7 7 7 7 7 presentation hemorrhage LENT Hemorrhage, ulceration Hemorrhage, ulceration ulceration Hemorrhage Hemorrhage Hemorrhage Hemorrhage Hemorrhage, Diarrhea, Pain Time to LENT diagnosis 17.5 15.5 14.5  $(mo)^*$ 10.5 17.5 17 13 27 12 X-ray, 6,000 cGy (200 cGy x 30 6,696 cGy X-ray, 5,000 cGy (200 cGy x 25 Brachytherapy, 3,239.5 cGy X-ray, 5,000 cGy 3,729 cGy X-ray, 5,000 cGy X-ray, 5,000 cGy X-ray, 5,000 cGy X-ray, 4,600 cGy X-ray, 5,000 cGy cGy (242.1 cGy x 19 fractions) 2,853 cGy X-ray, 4,599.9 Brachytherapy, 3,276 cGy (200 cGy x 25 fractions) Brachytherapy, 3,577 cGy Brachytherapy, Brachytherapy, (200 cGy x 25 (200 cGy x 25 Brachytherapy, Brachytherapy, (200 cGy x 25 (200 cGy x 23 Brachytherapy, Brachytherapy, (200 cGy x 25 3,503 cGy 3,808 cGy fractions) 3,500 cGy fractions) fractions) fractions) fractions) fractions) fractions) Cancer treatments Chemotherapy Irinotecan/ 1,478 mg (type/dose) Cisplatin/ 300 mg Cisplatin/ 360 mg 8 Š οŠ οÑ ž å Complementary TAH Complementary Surgery TAH(type) ν̈́ ž ž ž å ž ž ASCC/IIIb SCC/IIb SCC/IIb type/stage SCC/IIIb Cancer AC/IIb SCC/IIb SCC/IIb SCC/Ib2 SCC/IIb Tumor location cervix cervix cervix cervix cervix cervix cervix cervix cervix 7/21/1999 Uterine 4/23/1999 Uterine Uterine 10/1/1998 Uterine Uterine 11/1/1999 Uterine 3/16/2000 Uterine 6/12/2000 Uterine Uterine diagnosis 7/3/2000 Cancer 9/8/1999 7/4/2000 date Gender [T Ľ ſĽ, Ľ, Ľ, щ ſτ. Ľ, Ľ, PROC 020A PROC 024A PROC 021A PROC 023B PROC 025A PROC 026A PROC 027B PROC 022B PROC 028B Patient 

Yes	<sup>O</sup> N	oN O	Ň	°N	N	Yes	°Z	No	Yes	No (Continued)
N <sub>o</sub>	No.	No	No	Yes	Yes	No	No	No	No	Š
Yes	No	Yes	No	Yes	Š	Š	No	S <sub>O</sub>	No	Š
Never	Never	Never	Never	Past use	Never	Never	Never	Current use	Never	Never
61	2, 12 (Diet)	12 (Ferrous sulfate and diet)	2, 12 (Diet and metronidazole)	61	Ξ	2, 11	61	6	2, 12 (Diet)	12 (Ferrous Sulfate)
Hemorrhage, ulceration, Concurrent cystitis	Diarrhea, hemorrhage	Hemorrhage, ulceration	Hemorrhage, stricture	Diarrhea, pain, hemorrhage	Нетогћаде	Hemorrhage, ulceration, stricture	Hemorrhage	Hemorrhage	Hemorrhage	Hemorrhage, ulceration, stricture
4 3.	21	15.5	∞	∞	10	13.5	2.5	22	14.5	Ξ
X-ray, 4,600 cGy (200 cGy x 23 fractions) Brachytherapy,	3500 ccy X-ray, 5,200 cGy (400 cGy x 3 + 200 cGy x 20 fractions) Breachtherapy,	3,500 ccy X-ray, 4,800 cGy (300 cGy x 16 fractions) Brachytherapy,	3,500 cGy X-ray, 5,000 cGy (200 cGy x 25 fractions) Bractions,	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	3,500 cGy X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	4,352 cuy X-ray, 6,000 cGy (200 cGy x 30 fractions) Brachytherapy,	3.119 cGy X-ray, 5600 cGy (200 cGy x 28 fractions) Brachytherapy,	3,000 cOy X-ray, 6,000 cGy (200 cGy x 30	X-ray, 5,000 cGy (200 cGy x 25 fractions)  Brachytherapy,	3,528 cuy X-ray, 5,000 cGy (200 cGy x 19 + 400 cGy x 3 fractions) Brachytherapy, 3,500 cGy
Cisplatin/ 300 mg	Cisplatin/ 50 mg	°Z	°Z	°Z	Cisplatin/ 360 mg	°Z	Cisplatin/ 360 mg	No	Š	Š
N	°Z	N <sub>O</sub>	N	No	N	N <sub>o</sub>	oN	No	ТАН	ТАН
SCC/Ib2	SCC/Ib2	SCC/IIb	SCC/IIIb	SCC/IIIb	SCC/IIb	SCC/Ib2	SCC/IIIb	SCC/IIIb	SCC/IIa	ASCC/Ib2
Uterine cervix	Uterine cervix	Uterine cervix	Uterine cervix	Uterine cervix	Uterine cervix	Uterine cervix	9 Uterine cervix	9 Uterine cervix	Uterine cervix	Uterine cervix
11/1/1999 Uterine cervi	8/2/1999	5/15/2000 Uterine cervi	7/24/2000 Uterine cervi	6/29/2000 Uterine cervi	1/24/2000 Uterine cervi	5/2/2000	11/11/1999 Uterine cervi	10/20/1999 Uterine cervi	4/14/2000 Uterine cervi	2/12/2001 Uterine cervi
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PROC 029A	PROC 030B	PROC 031A	PROC 032B	PROC 033A	PROC 034A	PROC 035B	PROC 036B	PROC 037B	PROC 038B	PROC 039A

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						Table 9. Landin G	raore 5. ranem vemograpmes (commuted)	ninca)						
		Cancer				Cancer treatments		Time to		Previous				
Patient ID Ge	d Gender	diagnosis date	Tumor location	Cancer type/stage	Surgery (type)	Chemotherapy (type/dose)	di <i>RT/dosage</i>	diagnosis (mo)*	LENT	LENT treatment	Tobacco Diabetes use mellitus	Diabetes mellitus	Hypertension	Diabetes mellitus Hypertension Transfusions
PROC 040A	F 9/	6661/6/6	Uterine cervix	SCC/IIb	N <sub>O</sub>	°Z	X-ray, 5200 cGy (200 cGy x 20 + 400 cGy x 3 fractions) Brachytherapy,	41	Hemorrhage	2, 11	Never	No	oN O	Yes
PROC 041B	F 12	12/13/2000 Uterine cervi	Uterine cervix	SCC/IIIb	No	oN	3,792 cGy X-ray, 7,000 cGy (200 cGy x 35	9	Pain, hemorrhage	61	Never	No	No	No
PROC 042B	F 8/	8/25/1997 Uterine cervi	Uterine cervix	SCC/IIIb	N	o N	Hactrons) X-ray, 5,600 cGy (200 cGy x 28 fractions) Brachytherapy,	53.5	Hemorrhage, ulceration	6	Never	Yes	No	Yes
PROC 043A	F 2/	2/28/2001 Uterine cervi	Uterine cervix	SCC/IIb	No	Cisplatin/ 350 mg	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	13.5	Hemorrhage	12 (Diet)	Past use	Š	No	°
PROC 044A	F 2/	2/28/2001 Uterine	Uterine	SCC/IIb	No	Cisplatin/	Brachytherapy,	15.5	Hemorrhage	2	Past use	Š	No	Yes
PROC 045B	F 10	10/29/1999 Uterine cervi	Uterine cervix	SCC/IIIb	°Z	No oN	X-ray, 6,600 cGy (200 cGy x 33 fractions)  Brachytherapy,	44.5	Diarrhea, hemorrhage	2, 2,	Past use	Š	No	Yes
PROC 046A	F 7/	7/23/2001 Uterine cervi	Uterine cervix	SCC/IIb	°Z	Cisplatin/ 360 mg	3,500 cGy X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	10.5	Diarrhea, pain, hemorrhage, ulceration, stricture	m	Past use	Š	N <sub>O</sub>	Ñ
PROC 047A	F 4/	4/18/2000 Uterine cervi	Uterine cervix	ASCC/Ib1	°Z	°Z	2019 COY X-ray, 7,000 cGy (200 cGy x 25 + 200 cGy x 10 fractions)	26.5	Bleeding, metabolic disorder	12 (Diet)	Never	Yes	N <sub>O</sub>	Yes
PROC 048B	M 51	10/7/2000 Prostate	Prostate	AC	°Z	°Z	X-ray, 6,840 cGy (180 cGy x 38 fractions)	17	Diarrhea, pain, hemorrhage, fistula, edematous	2, 3	Current use	Š	Yes	No
PROC 049A	F 6/	6/1/2001	Uterine cervix	SCC/IIIb	<sup>o</sup> Z	Cisplatin/ 200 mg	X-ray, 6,000 cGy (200 cGy x 30 fractions) Brachytherapy,	13	Diarrhea, pain, hemorrhage, ulceration,	C1	Never	No	No	Yes

Yes	Yes	Yes	N O	No	° X	°Z	°Z	Yes	Yes	No (Continued)
No	No	No	Yes	No	N <sub>O</sub>	No	No	No	Yes	Š
No	N <sub>o</sub>	N <sub>o</sub>	No	No	Š	S <sub>o</sub>	Š	No	No	Š
Never	Never	Past use	Never	Never	Never	Never	Never	Never	Never	Never
2, 12 (Diet)	-	71	6	12 (Diet)	<i>C</i> I	-	12 (Diet)	12 (Steroid use)	12 (Steroid use)	2, 12 (Diet, steroid enema)
Hemorrhage	Pain, ulceration	Diarrhea, pain, hemorrhage, ulceration	Hemorrhage	Hematuria	Hemorrhage	Hemorrhage	Hemorrhage	Hemorrhage	Hemorrhage	Cramping, pain, hemorrhage
51.5	12.5	18.5	20		21	16.5	28	12.5	18	16.5
X-ray, 6,000 cGy (200 cGy x 30 fractions) Brachytherapy,	5,495 ccyy X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	Z,936 cCy X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	5,331 CUY X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	2,6/0 ccty X-ray, 7600 cGy (200 cGy x 38 fractions) Brachytherapy,	Z,480 cCy X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	5,27.8 ctty X-ray, 5,100 cGy (300 cGy x 17 fractions) Brachytherapy,	5,400 cGy X-ray, 4,600 cGy (200 cGy x 23 fractions) Brachytherapy,	5,450 cdy X-ray, 5,000 cdy (200 cdy x 25 fractions) Brachytherapy,	2,762 CGy X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	3,777 cuy X-ray, 5,000 cGy (200 cGy x 25 fractions) Bracchytherapy, 3,022 cGy
°Z	°Z	Cisplatin/ 350 mg	°Z	°Z	°Z	o N	o N	o N	o N	Cisplatin/ 240 mg
N	Š.	N <sub>o</sub>	ТАН	N O	N <sub>o</sub>	°N	°Z	°N	N <sub>o</sub>	ŝ
SCC/IIa	SCC/IIb	SCC/IIa	SCC/IIb	Cancer epidermoid/IIb	SCC/IIIb	SCC/IIIb	SCC/IIa	SCC/Ib2	SCC/Ib1	SCC/IIIb
6/23/1997 Uterine cervix	4/30/2001 Uterine cervix	1 Uterine cervix	4/14/2000 Uterine cervix	12/3/2002 Uterine cervix	3/14/2001 Uterine cervix	0 Uterine cervix	4 Uterine cervix	11/8/2000 Uterine cervix	11/14/2000 Uterine cervix	11/30/2000 Uterine cervix
6/23/19	4/30/20	5/7/2001	4/14/20	12/3/20	3/14/20	3/3/2000	9/3/1984	11/8/20	11/14/2	11/30/2
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PROC 050B	PROC 051B	PROC 052A	PROC 053B	PROC 054A	PROC 055A	PROC 056B	PROC 057A	PROC 058A	PROC 059B	PROC 060B

(Continued)

							`						
	(		,		Cancer treatments		Time to						
Patient ID Ge	Cancer diagnosis Gender date	er sis Tumor location	Cancer type/stage	Surgery (type)	Chemotherapy (type/dose)	CT/dosage	diagnosis $(mo)^*$	s LENT presentation	Previous LENT treatment <sup>†</sup>	Tobacco	Diabetes mellitus	Hypertension	Diabetes mellitus Hypertension Transfusions
PROC 061A	F 1/3/2001	1 Uterine cervix	SCC/IIIb	No	N <sub>o</sub>	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	16	Diarrhea, hemorrhage, cramping	2	Never	No	No	No
PROC 062B	F 11/26/19	11/26/1998 Uterine cervix	SCC/Ib2	No	°N	4,229 cO.y X-ray, 6,000 cGy (200 cGy x 30 fractions) Brachytherapy,	10.5	Hemourhage	6	Never	Š	Yes	Yes
PROC 063A	F 12/13/2	12/13/2000 Uterine cervix	SCC/IIIb	°Z	°Z	5,502 cuy X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	19.5	Hemorrhage	6	Never	Š	No	Yes
PROC 064B	F 9/24/19	9/24/1999 Uterine cervix	SCC/IIb	No	Cisplatin/ 420 mg	X-ray, 5600 cGy (200 cGy x 28 fractions) Brachytherapy,	41	Diarrhea, pain, hemorrhage	7	Never	No	Š	°Z
PROC 065A	F 11/6/20	11/6/2001 Uterine cervix	SCC/IIb	Radical hysterectomy	Cisplatin/ 420 mg	X-ray, 5,000 cGy (200 cGy x 25 fractions)	8.5	Hemorrhage, edematous wall	6	Never	No	No	Yes
PROC 066A	M 5/1/2001	1 Prostate	AC	No	°Z	X-ray, 6840 cGy (180 cGy x 38 fractions)	19	Hemorrhage, ulceration, Wall changes (Mucosal	62	Current use	Š	Yes	N <sub>o</sub>
PROC 067B	F 7/16/20	7/16/2001 Uterine cervix	SCC/IIb	No	Cisplatin/ 390 mg	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy, 2,500 cG.	6	Pain, hemorrhage, ulceration, stricture	2	Never	No	N <sub>o</sub>	°Z
PROC 068B	F 11/21/2	11/21/2001 Uterine cervix	SCC/IIIb	No	Carboplatin/ 450 mg	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	16	Hemorrhage, ulceration, edematous wall changes	6	Never	Yes	Š	Yes
PROC 069A	F 12/11/2	12/11/2001 Uterine cervix	SCC/IIb	No	Cisplatin/ 300 mg	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy, 3 321 cGy	8.5	Diarrhea, hemorrhage	6	Never	No	Yes	Yes
PROC 070B	F 9/20/20	9/20/2002 Uterine cervix	SCC/IIIb	Š	°N	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	13.5	Constipation, hemorrhage, ulceration	6	Never	No	Yes	°Z

<b>%</b>	Yes	Š	No	No	Yes	No	Yes	Yes	No (Continued)
No	No	Yes	S <sub>O</sub>	No	No	No	No	N <sub>0</sub>	No
Š	Yes	N <sub>O</sub>	°Z	No	Š	Š	Š	<sup>o</sup> Z	No
Never	Never	Past use	Never	Never	Never	Never	Never	Never	Never
61	12 (Diet)	N A	2	2, 12 (Dilatation)	61	2, 12 (Metronidazole)	2, 12 (Diet)	2, 3, 5, 12 (Steroid enema)	2, 12 (Steroid enema)
Нетотнаде	Hemorrhage	Pain, hemorrhage, stricture, wall changes (edematous, electronic)	Hemorrhage	Hemorrhage, ulceration, stricture	Hemorrhage	Constipation, hemorrhage ulceration	Hemorrhage	Diarrhea, Constipation, pain, Hemorrhage, wall changes (edematous, mucosal thickening), other (hyperemia,	erosions) Hemorrhage
13.5	3.5	4	17.5	16.5	16	17.5	9.5	10	119
X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy, 3,200 cGy	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	X-ray, 5040 cGy (180 cGy x 28 fractions)	X-ray, 5,312 cGy (180 cGy x 29 fractions) Brachyherapy, 6,804 cGy	X-ray,5,000 cGy (200 cGy x 25 fractions) Brachytherapy, 3,085 cGv	X-ray, 6,000 cGy (200 cGy x 30	Tractions)  X-ray, 5400 cGy (200 cGy x 27 fractions)  Brachytherapy, 3.486, cGs.	S,450 CGy X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	3,701 COy X-701 COy Z-00 cGy x 25 fractions) Brachytherapy, 2,750 cGy	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy, 3,077 cGy
Cisplatin/ 350 mg	Š.	FU (400 mg) + FA (20 mg)	PVC (before RT, platinum 150 mg, after RT vincristine 400 mg 400 mg with platinum 40 mg vincristine 400 mg vincristine 400 mg vincristine 40 mg vincristine	Cisplatin/300 mg	Cisplatin/300 mg	Cisplatin/ 450 mg + Cyclophosphamide/ 4,500 mg	<sup>O</sup> N	Cisplatin/350 mg	Ň
°N	N <sub>o</sub>	Low anterior resection + end-to-end anastomoses	Hysterectomy + BSO	°N	No	TAH	Ŝ	Ŝ	Š
SCC/III	SCC/IIb	AC	Other (glassy cells)/IIb	SCC/IIb	ASCC/IIIb	AC	SCC/IIIb	SCC/IIIa	SCC/IIIb
11/21/2001 Uterine cervix	6/20/2001 Uterine cervix	10/28/1998 Rectum	4/5/2002 Uterine cervix	7/26/2000 Uterine cervix	4/18/2002 Uterine cervix	5/3/2002 Uterine corpus	6/11/2000 Uterine cervix	3/3/2003 Uterine cervix	5/20/2002 Uterine cervix
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PROC 071A	PROC 072B	PROC 073B	PROC 074B	PROC 075A	PROC 076A	PROC 077A	PROC 078B	PROC 079A	PROC 080B

Table 3. Patient demographics (continued)

						Concer transferants	Table 3. Fauefit demographics (continuea)	Time to						
	Ü	Cancer				Califor deadillonis		LENT		Previous				
Patient ID Ge	diag Gender da	diagnosis date	Tumor location	Cancer type/stage	Surgery (type)	Chemotherapy (type/dose)	d <i>RT/dosage</i>	diagnosis (mo)*	LENT presentation	LENT treatment <sup>†</sup>	Tobacco Diabetes use mellitus	Diabetes mellitus	Diabetes mellitus Hypertension Transfusions	Fransfusions
PROC 081A	F 5/10/	5/10/2001 U	Uterine corpus	AC	Hydrothermal ablation	%	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	41	Hemorrhage	2	Never	No	No	No
PROC 082A	F 1/7/2002		Uterine corpus	AC	TAH	Š.	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	22	Hemorrhage	61	Never	Yes	Yes	No
PROC 083B	F 9/2/2002		Uterine cervix	SCC/IIIb	Š	Gemzar/ 2700 mg	X-ray, 5,600 cGy (200 cGy x 28 fractions) Brachytherapy,	11.5	Hemorrhage	61	Never	Š	Š	Yes
PROC 084B	F 2/15/	2/15/2003 Uterine corpu	rerine	AC	ТАН	Ž	X-152 COy X-ray, 5040 cGy (180 cGy x 28 fractions) Brachytherapy, 1,800 cGy	12.5	Cramping, constipation, pain, hemorrhage, ulceration, endarteritis, wall changes (Ademotive)	3, 12 (Coagulation by adrenaline injection and heater probe)	Never	<sup>o</sup> Z	Ŝ	Yes
PROC 085A	F 1/25/	1/25/2002 Uterine cervi	ferine cervix	SCC/IIb	°Z	No	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy, 2,217,25.	19.5	(coonantage Hemorrhage	7	Never	Š	Š	°N
PROC 086B	F 1/21/	1/21/2003 Uterine corpu	terine corpus	AC	TAH + BSO, node sampling	N <sub>o</sub>	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy, 5,000 cG.	10.5	Pain, hemorrhage, ulceration	NA	Never	Š	No	Yes
PROC 087B	F 7/2/2002		Uterine corpus	AC	TAH	No	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	∞	Hemorrhage	7	Never	No	N <sub>o</sub>	No
PROC 088A	F 7/5/2002		Uterine cervix	SCC/IIa	°Z	Carboplatin/ 200 mg	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	8.5	Hemorrhage	6	Never	Yes	Yes	Yes
PROC 089B	F 5/24/	5/24/2002 Uterine cervi	fterine cervix	SCC/IIb	°Z	No	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachyherapy,	19.5	Hemorrhage	2	Never	N	No	N <sub>o</sub>

N	Yes	No	°Z	Š	Yes	%	No	°Z	Ñ	Yes	No (Continued)
No	Š	Yes	Ö	Š	No	o Z	Yes	No	No	N <sub>o</sub>	N
Š	No	oN O	No	No	No	No	$ m N_0$	Š	No	No	Yes
Current use	Never	Past use	Past use	Past use	Never	Never	Past use	Past use	Never	Never	Current use
12 (Diet)	'n	2,5	3, 4, 5	61	1,2	2, 12 (Diet)	2,5	, , , , , , , , , , , , , , , , , , ,	2	2	2, 3, 5
Cramping, pain, stricture, Perforation	Pain, hemorrhage, ulceration	Pain, hemorrhage, ulceration	Diarrhea, vomiting, pain, Cramping, hemorrhage	Hemorrhage, ulceration	Pain, hemorrhage, wall changes	(edematous) Hemorrhage	Constipation, pain, hemorrhage,	charterius Cramping, pain, hemorrhage, hypocellularity, hypovascularity, wall changes	(edematous) Diarrhea, pain, hemorrhage	Hemorrhage, ulceration	Diarrhea, cramping, pain
41	29.5	Ξ	106	8.5	Ξ	27	61	10.5	91	10.5	20.5
X-ray, 5,500 cGy (183.33cGy x 30 fractions) Brachytherapy,	2,000 cGy Co60 (pendulum), 6,750 cGy (250 cGy x 27	X-ray, 7,200 cGy (200 cGy x 36	X-ray, 4,500 cGy (1.8cGy x 25 fractions)  Brachytherapy,	6,000 cGy X-ray, 5,000 cGy (400 cGy x 3 + 200 cGy x 19 fractions) Brachytherapy,	3,147 cuy X-ray, 6,400 cGy (200 cGy x 32	racuons) X-ray, 5,600 cGy (200 cGy x 28 fractions) Brachytherapy,	3,100 cGy X-ray, 7,400 cGy (200 cGy x 37	rractions) X-ray, 7,200 cGy (180 cGy x 40 fractions)	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	2,990 cuy X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	5,2-1 cuy X-ray, 4,230 cGy (176,2cGy x 24 fractions) Brachytherapy, 4,500 cGy
Cisplatinl390 mg	%	oN.	Cisplatin Adriamycin (dose unknown)	Cisplatin/280 mg	oN	°Z	No	Neoadjuvant hormonal therapy	Cisplatin/ 170 mg	Cisplatin/ 360 mg	Cisplatin/ 190 mg
Cone biopsy	No	No	TAH + BSO, lymphadenectomy	<sup>O</sup> N	TAH + BSO	Ň	No	N <sub>O</sub>	No	Ň	Š.
SCC/IIIa	SCC/IIIb	AC	Carcinosarcoma (mixed malignant mullerian tumor)	SCC/IIb	Adenosarcoma	SCC/IIIb	AC	AC	SCC/IIb	AC	SCC/IIb
Uterine cervix	Uterine cervix	3/28/2003 Prostate	Uterine	Uterine cervix	Uterine corpus	Uterine cervix	5/28/1999 Prostate	Prostate	Uterine cervix	Uterine cervix	Uterine cervix
5/20/2001 Uterine	3/26/2001 Uterine cervi	3/28/2003	10/1/1990 Uterine corp	4/4/2003	4/24/2002 Uterine corp	1/7/2000	5/28/1999	2/13/2002 Prostate	12/6/2000 Uterine cervi	11/7/2002 Uterine cervi	9/3/2002
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PROC 090A	PROC 091B	PROC 092A	PROC 093B	PROC 094A	PROC 095A	PROC 096B	PROC 097B	PROC 098A	PROC 099A	PROC 100A	PROC 101A

(Continued)

Table 3. Patient demographics (continued)

	Diabetes mellitus Hypertension Transfusions	No	Yes	N <sub>o</sub>	Š	Yes	Yes	Yes	Yes
	Hypertension	No	Š	Yes	Ŷ	Yes	Š	No No	Yes
	Diabetes mellitus	S <sub>o</sub>	No	Ň	Š	Š	o N	No	Š
	Tobacco	Current use	Past use	Never	Current use	Past use	Never	Never	Never
	Previous LENT treatment <sup>†</sup>	1, 2, 3	2	2, 5, 7	1, 2	A A	2, 12 (Diet)	71	1, 2
	LENT presentation	Constipation, pain, hemorrhage,	Cramping, pain, hemorrhage	Diarrhea, cramping, ulceration, stricture, Endarteritis, hypocellularity, hymorogenelarity, hymorogene	Diarrhea, cramping, pain, hemorrhage, wall changes (edematous, fibrous), other (telangeictasia, recional arrowhy)	Vomiting, Constituting, hemorrhage, ulceration, stricture, wall changes (defenatous, financia)	Hemorrhage	Hemorrhage	Vomiting, cramping, pain, Constipation, hemorrhage, ulceration, wall changes
Time to	LENT diagnosis $(mo)^*$	18	17	∞	13	6	15.5	12.5	28.5
	RT/dosage	X-ray, 7,000 cGy (200 cGy x 35	X-ray, 6,750 cGy (250 cGy x 27 fractions)	Tractions)  X-ray, 5,250 cGy (175 cGy x 30 fractions)	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy, 2,000 cGy	X-ray, 5,000 cGy (200 cGy x 25 fractions)	X-ray, 5,800 cGy (200 cGy x 25 fractions + 800 cGy) Brachytherapy, 2,950 cGv	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy, 2,660 cGy	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy, 2,200 cGy
Cancer treatments	Chemotherapy (type/dose)	No	Cisplatin/ 55 mg	Ö	Ž	Ž	Ö	%	Ö
	Surgery (type)	No	No	TAH, pelvic node dissection and omental biopsy	TAH + BSO	Radical hysterectomy + bilateral iliac lymph node dissection	N <sub>O</sub>	°Z	TAH + BSO
	Cancer type/stage	AC	SCC/IIIb	AC	SCC/lb1	AC	SCC/IIa	AC	AC
	Tumor location	Prostate	Uterine cervix	Uterine cervix	Uterine cervix	12/21/2003 Endometrium	Uterine cervix	Uterine corpus	Uterine corpus
	Cancer diagnosis date	1/28/2003	4/26/2003	1/1/2000	9/1/2000	2/21/2003	2/19/2003 Uterine cervi	2/8/2002	6/10/2002 Uterine corpu
	Gender	M 1,	Ŧ ,4	F 1	년 6	г.	F 2	F 2	F Q
	Patient ID (	PROC 102A	PROC 103B	PROC 104B	PROC 105B	PROC 106A	PROC 107B	PROC 108A	PROC 109A

N <sub>O</sub>	Yes	°Z	°N	Yes	Yes	No	No	°Z	°Z	No	No (Continued)
N <sub>o</sub>	No	o N	Yes	o N	o O	No	No	o N	o N	N <sub>o</sub>	°Z
No	No	N	No	No	No	No	No	Yes	N	No	°Z
Never	Never	Never	Never	Never	Never	Never	Never	Never	Never	Never	Never
6	6	0	0	2, 3, 5	2, 5	NA	3,5,7,9	6	0	NA	5, 12 (Analgesic: morphine)
Hemorrhage	Hemorrhage	Hemorrhage	Hemorrhage	Diarrhea, pain, hemorrhage	Diarrhea, pain, hemorrhage	Hemorrhage, wall changes	Diarrhea, cramping, pain, Constipation,	hemorrhage Constipation, pain, Hemorrhage, ulceration, Wall changes (Pale, edematous,	riototic) Diarrhea, hemorrhage	Diarrhea, cramping, pain	Diarrhea, cramping, pain, constipation, ulceration, stricture, wall changes (edematous, mucosal thickening)
21.5	12.5	17.5	10.5	17	16	23.5	126	17	10.5	NA	17.5
X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	3,303 CGy X-ray, 7,000 cGy (200 cGy x 35	X-ray, 5,000 cGy (200 cGy x 25 fractions)  Brachytherapy,	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	2,934 cOy X-ray, 6,750 cGy (250 cGy x 27	X-ray, 5,250 cGy (250 cGy x 21	Hactions) X-ray, 7,000 cGy (200 cGy x 35 fractions)	X-ray, dosage unknown	X-ray, 6,480 cGy (180 cGy x 36 fractions)	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	Z,700 CGy X-ray, 6600 cGy (200 cGy x 33	ractions) X-ray, 5199cGy (173.3cGy x 30 fractions) Brachytherapy, 1,800 cGy
Cisplatin/55 mg & Gemzar/175 mg	Carboplatin/ 350 mg	Ŝ	Cisplatin/ 300 mg	No	No	Hormonal therapy	N <sub>o</sub>	Casodex/50 mg lucrin depot 3 M/ 11.25 mg	Ŝ	No	Cisplatin/330 mg
Radical hysterectomy	Radical hysterectomy	N <sub>o</sub>	No	No	No	No	Resection with colostomy	Transurethral resection	N <sub>O</sub>	No	TAH
SCC/IIb	AC	SCC/IIb	SCC/IIIa	SCC/IIIb	SCC/IIIb	AC	AC	AC	SCC/Ib1	AC	SCC/Ib2
Uterine cervix	Uterine cervix	Uterine cervix	Uterine cervix	Uterine cervix	Uterine cervix	Prostate	Colon	Prostate	Uterine cervix	Prostate	Uterine cervix
2/13/2002 Uterine	2/20/2003 Uterine cervi	8/8/2003	5/14/2003 Uterine cervi	1/16/2003 Uterine cervi	4/14/2003	5/1/2002	8/1/1987	5/1/2003	1/24/2004 Uterine cervi	NA	6/6/2003
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PROC 110A	PROC 111B	PROC 112B	PROC 113B	PROC 114B	PROC 115A	PROC 116A	PROC 117A	PROC 118B	PROC 119B	PROC 120A	PROC 121B

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	,			'		Cancer treatments		Time to						
Patient ID G	diz Gender	Cancer diagnosis date	Tumor location	Cancer type/stage	Surgery (type)	Chemotherapy (type/dose)	C RT/dosage	LENT diagnosis $(mo)^*$	LENT presentation	Previous LENT treatment <sup>†</sup>	Tobacco	Diabetes mellitus H	Diabetes mellitus Hypertension Transfusions	ransfusions
PROC 122A	F 1/1	1/14/2002 1	Uterine cervix	SCC/IIIb	No	Carboplatin/ 1505 mg	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	18	Diarrhea, hemorrhage	2	Never	No	No	No
PROC 123A	F 3/1.	3/14/2002 Uterine cervi	Uterine cervix	SCC/IIb	Ν̈	S.	2,860 cGy X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	18.5	Diarrhea, pain, hemorrhage, ulceration, wall changes (fibrotic)	1, 2, 3, 5	Never	No	Yes	Yes
PROC 124B	F 9/2	9/22/2003 Uterine cervi	Uterine cervix	SCC/IIb	<sup>Q</sup> Z	Carboplatin/ 600 mg	2,730 cGy X-ray,5,000 cGy (400 cGy x 3 + 200 cGy x 19 fractions) Brachytherapy, 3,311 cGy	7.5	Diarrhea, hemorrhage	71	Never	No	°Z	N
PROC 125B	F 1/2	1/27/1987 Uterine cervi	Uterine cervix	SCC/IIIb	Staging laparotomy, debulking of enlarged nodes in pelvis and transposition of left every	°Z	X-ray, 5220 cGy (180 cGy x 29 fractions) Brachytherapy, 2,100 cGy	155	Diarrhea, cramping, pain, wall changes (edematous)	1,2,5	Current	°Z	°Z	N O
PROC 126B	F 7/1	7/15/2003 Uterine cervi	Uterine cervix	SCC/IIa	TAH + BSO	Cisplatin/ 120 mg	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	12	Cramping, pain, hemorrhage	2, 3, 5	Never	No	Yes	Yes
PROC 127A	F 1/1	1/1/2004	Uterine cervix	SCC/IIIb	N <sub>o</sub>	No	X-ray, 6,750 cGy (250 cGy x 27	6	Pain, hemorrhage	2, 3, 5	Never	Yes	Yes	Yes
PROC 128A	F 8/1	8/1/2003 [	Uterine cervix	SCC/Ib2	N <sub>O</sub>	Cisplatin/ 330 mg	X-ray, 5,600 cGy (200 cGy x 28 fractions) Brachytherapy, 2020 cG.	4	Constipation, hemorrhage	61	Never	No	No	N <sub>o</sub>
PROC 129B	F 10/	10/8/2003 Uterine cervi	Uterine cervix	ASCC/IIIb	Ö	Cisplatin/ 330 mg	X-ray, 6,000 cGy (200 cGy x 30 fractions)  Brachytherapy, 7879 cGy	11.5	Нетопнаде	71	Never	No	Š	N <sub>O</sub>
PROC 130A	F 4/4	4/4/2004 1	Uterine cervix	SCC/IIb	Š	Cisplatin/ 240 mg	X-ray, 5,000 cGy (400 cGy x 3 + 200 cGy x 19 fractions) Brachytherapy,	10	Pain, hemorrhage, ulceration	2	Never	°Z	S Z	Yes

Yes	°N	Yes	No	Yes	Ö	No	No	Yes	°Z	No	Yes	Yes (Continued)
No	No	N <sub>o</sub>	°Z	o N	Yes	Yes	No	°Z	Ö	°Z	°Z	Yes
No	No	No	No	No	No	Yes	No	N <sub>o</sub>	Yes	N <sub>o</sub>	No	No
Never	Never	Never	Never	Never	Never	Never	Past use	Never	Never	Never	Never	Never
2, 5, 10	2, 3, 5	61	61	61	7	2,5	4,5	61	7	61	<b>C</b> 1	, 2, 5,
Cramping, pain, hemorrhage	Diarrhea, cramping, pain, hypovascularity, wall changes (fibrotic, mucosal thickenine)	Pain, hemorrhage	Wall changes (edematous)	Pain, hemorrhage	Wall changes (edematous) other (telangiectasia)	Diarrhea, cramping, pain	Cramping, pain, hemorrhage	Hemorrhage	Hemorrhage, wall changes (edematous)	Hemorrhage, endarteritis, wall changes (edematous)	Hemorrhage	Pain, hemorrhage
12.5	16.5	15	19	1	15.5	4.5	49.5	41	15.5	12.5	19	17.5
X-ray, 6,750 cGy (250 cGy x 27 fractions)	X-ray, 4000 cGy (160 cGy x 25 fractions) Brachytherapy dosage unknown	X-ray, 5,400 cGy (200 cGy x 27	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy, 2156 cGv	X-ray, 6,750 cGy (250 cGy x 27 fractions)	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy, 3 200 cGv	X-ray, 6,300 cGy (210 cGy x 30 fractions)	X-ray, 6800 cGy (200 cGy x 34 fractions)	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy, 3,430 cGy.	X-ray, 5,600 cGy (200 cGy x 28 fractions) Brachytherapy, 4 841 cGy	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy, 3 508 cGy	X-ray, 6,000 cGy (200 cGy × 30 fractions) Brachytherapy, 7,65, cGv	X-ray, 6,750 cGy (250 cGy x 27 fractions)
Cisplatin/ 175 mg	No	No	Carboplatin/ 900 mg	Cisplatin/90 mg	ON	No	Hormonal therapy	Cisplatin/ 350 mg	Cisplatin/50 mg	Cisplatin/70 mg	Cisplatin/350 mg	Cisplatin/136 mg
No	ТАН	No	No	No	°N	No	No	No	°	No	oN	No
SCC/IIIb	SCC/IIa	SCC/IIIb	SCC/IIb	SCC/IIIb	SCC/?	AC	AC	SCC/Ib2	AC	SCC/IIb	Squamous transitional papilar cell carcinoma	SCC/IIIb
3/25/2002 Uterine cervix	2/7/1993 Uterine cervix	11/19/2003 Uterine cervix	7/28/2003 Uterine cervix	9/16/2003 Uterine cervix	3/8/2004 Uterine cervix	9/1/1999 Prostate	12/15/2000 Prostate	3/30/2004 Uterine cervix	12/16/2003 Uterine cervix	3/2/2004 Uterine cervix	4/29/2002 Uterine cervix	12/8/2003 Uterine cervix
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PROC 131A	PROC 132B	PROC 133B	PROC 134A	PROC 135B	PROC 136A	PROC 137B	PROC 138A	PROC 139B	PROC 140A	PROC 141A	PROC 142A	PROC 143B

Table 3. Patient demographics (continued)

					Cancer treatments		Time to						
Gender	Cancer diagnosis date	Tumor location	Cancer type/stage	Surgery (type)	Chemotherapy (type/dose)	d <i>RT/dosage</i>	LENT diagnosis (mo)*	LENT presentation	Previous LENT treatment <sup>†</sup>	Tobacco Diabetes use mellitus	Diabetes mellitus E	Diabetes mellitus Hypertension Transfusions	Fransfusions
	10/14/2003 Prostate	) Prostate	AC	TURP	No	X-ray, 7,000 cGy (200 cGy x 35 fractions)	11.5	Diarrhea, pain, hemorrhage, Hypocellularity, hypovascularity, wall changes	1, 2	Never	No	oN	No
	5/6/2004	Uterine cervix	SCC/IIb	Š	Cisplatin1300 mg + gemcetabinel 1000 mg	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	8.5	(pale) Hemorrhage	2	Never	Š	°Z	N <sub>o</sub>
	M 10/17/2003 Prostate	Prostate	AC	No	No	check dosage X-ray, 4,500 cGy (180 cGy x 25 fractions)	6	Diarrhea, cramping, pain, hemorrhage, wall changes	61	Past use	No	Yes	N <sub>o</sub>
	3/10/2003 Rectum	Rectum	AC	Low anterior resection	5-FU/15 g	X-ray, 5,040 cGy (180 cGy x 28 fractions)	9.5	(edematous) Diarrhea, constipation, pain, hemorrhage,	NA	Never	No	Yes	Yes
	3/26/2004	3/26/2004 Uterine corpus	Mix mesodermal tumor (carcinosarcoma)	TAH	No	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy, 2 510 cG.	13.5	Diarrhea, hemorrhage	61	Never	No	o <sub>N</sub>	No
	10/20/2003	10/20/2003 Uterine cervix	SCC/IIb	Ŝ	Cisplatin/300 mg	X-ray, 5,000 cGy (200 cGy x 25 fractions) Brachytherapy,	23.5	Hemorrhage, endarteritis, wall changes (edematous)	61	Never	Š	o N	Š
	12/19/2000 Prostate	) Prostate	AC	N <sub>o</sub>	Hormonal therapy	7-702 CO3 X-ray, 6,600 cGy (200 cGy x 33 fractions)	1	Diarrhea, cramping, hemorrhage, wall changes (pale, fibrotic, mucosal thickening)	3, 11	Never	No	Yes	N <sub>o</sub>

Abbreviations: RT = radiotherapy; LENT = late effects normal tissue; SCC = squamous cell carcinoma; AC = adenocarcinoma; BSO = bilateral salpingo-oophorectomy; TAH = total abdominal hysterectomy; ASCC = Adenosquamous cell carcinoma; FU = Fluorouracil; FA = Folinic acid; PVC = portal vein chemotherapy; TURP = transurethral resection of prostate. \* Rounded to nearest month.

† Previous LENT treatment: 1 = antibiotics; 2 = anti-inflammatory agent; 3 = antispasmodic agents; 4 = anticholinergic agents; 5 = antidiarrheal agents; 6 = intestinal bypass; 7 = intestinal

resection; 8 = fistula repair; 9 = colostomy; 10 = ileostomy; 11 = fulguration; 12 = other.