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# Rectum separation in patients with cervical cancer for treatment planning in primary chemo-radiation

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# Abstract

**Purpose:** To proof feasibility of hydrogel application in patients with advanced cervical cancer undergoing chemo-radiation in order to reduce rectal toxicity from external beam radiation as well as brachytherapy.

**Material and methods:** Under transrectal sonographic guidance five patients with proven cervical cancer underwent hydro gel (20 cc) instillation into the tip of rectovaginal septum adherent to posterior part of the visible cervical tumor. Five days after this procedure all patients underwent T2 weighted transversal and sagittal MRI for brachytherapy planning. MRI protocol included T2 weighted fast spin echo (FSE) imaging in sagittal, coronal and para-axial orientation using an 1.5 Tesla MRI. Separation of anterior rectal wall and cervix was documented.

**Results:** Hydrogel application was uneventful in all patients and no toxicity was reported. Separation ranged from 7 to 26 mm in width (median 10 mm). The length of the separation varied between 18 and 38 mm (median 32 mm). In all patients displacement was seen in the posterior vaginal fornix, and/or at the deepest part of uterine cervix depending on the extension of the cul-de-sac in correlation to the posterior wall of the uterus. In patients with bulky tumor and/or deep (vaginal) extend of peritoneal cavity tumour was seen mainly cranial from the rectovaginal space and therefore above the hydrogel application. Only in the extra-peritoneal (lower) part of the cervix a good separation could be achieved between the rectum and cervix.

**Conclusion:** Hydrgel instillation in patients with cervial cancer undergoing chemoradiation is safe and feasible. Because of the loose tissue of the cul-de-sac and its intra- and extraperitoneal part, hydrogel instillation of 20 cc did not result in a sufficient separation of the cervix from anterior wall.

# Introduction

Primary chemoradiation (RCTX) is the treatment of choice in patients with locally advanced and/or lymph node positive cervical carcinoma [1]. Paradigm shift from radiation to RCTX lead to an improvement with regard to local control as well as progression free and overall survival [2]. Analysis of patterns of recurrence showed that locoregional control remains critical [3]. Nevertheless RCTX can be associated with considerable acute and late gastrointestinal (GI) toxicity [1,4]. Reported grade 3 and 4 late GI toxicity is in the range of 4%–40% depending on target volumes and radiation

techniques used [5-8]. However, possible improvement of local control due to dose escalation has to take GI and genitourinary (GU) toxicity into account.

Compared with 3D era IMRT based techniques could demonstrate lower rates of high grade toxicity [9-12]. Efforts have been made to better target definition for external beam radiation and dose escalation [13-15], dose prescription and application of brachytherapy, as well as simultaneous consideration of therapy related toxicity. Some proposals for dose sparing to the small bowel like treatment in prone position or use of bowel displacement systems did not gain acceptance in clinical routine [16,17].

Although external beam radiation (EBRT) as well as brachytherapy (BT) may lead to rectal complications only few data available concerning rectal toxicity in the



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treatment of cervical cancer patients. Their comparability is limited because of different treatment regimens, doses of EBRT and BT, dose prescriptions, application techniques and toxicity scoring systems. In cervical cancer patients cumulative overall and  $\geq 2$  grade rectal toxicity has been recorded in 12%–19% [18,19].

Injection of human collagen into the space between prostate and rectum in prostate cancer patients have shown decline of rectal dose by 25%–50% [20-24]. Therefore this new technique was thought to be an interesting option even for patients with cervical carcinoma undergoing EBRT and BT. A separation of at least 10–15 mm would be sufficient to achieve 80% rectal dose reduction [20]. Present study was initiated to proof feasibility of hydrogel application in patients with advanced cervical cancer in order to reduce rectal toxicity. To our best knowledge the present study is the first reporting of hydrogel application in patients with cervical cancer.

# **Material and methods**

This pilot study comprises five patients with proven cervical cancer (age 31-69 years, BMI 20-49 kg/m<sup>2</sup>). FIGO stages were IB1 pN1 in one, IIB in two and IIIB in two patients, respectively. All patients underwent primary RCTX as previously described [25]. During third week of EBRT patients underwent implantation of Smit-Sleeve® applicator under general anaesthesia (Varian, Palo Alto, CA) for brachytherapy. During this procedure hydro gel installation was also performed. Written informed consent was given by all patients. The present pilot study was IRB approved. Patient was placed in with adducted legs placed on leg holders. The procedure started by filling the bladder with 300 cc isotonic saline solution. Under abdominal sonography external cervical os was identified, dilated and Smit Sleeve was inserted. After fixation of the applicator with non resorbable sutures graspers were placed at 05.00 and 07.00 o'clock at vaginal introitus. By pulling the graspers downwards rectovaginal septum was brought under tension before gel application (Figure 1). Now a vaginal sonography probe was inserted into the rectum. Under transrectal sonographic guide hydrogel applicator (SpaceOAR®, Augmenix, Waltham, MA) was placed up to the tip of rectovaginal septum adherent to posterior part of the visible cervical tumour. Injection was done bringing the two components of the gel (20 cc) together (Figure 2).

Five days after this procedure all patients underwent T2 weighted transversal and sagittal MRI for brachytherapy planning. MRI protocol included T2 weighted fast spin echo (FSE) imaging in sagittal, coronal and paraaxial orientation using an 1.5 Tesla GE Signa Excite (GE Healthcare, Fairfield, CO, USA).



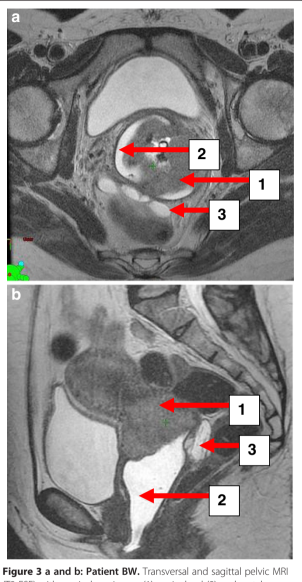
**Figure 1** Grasters to the vaginal introitus in order to bring the rectovaginal septum under tension before application.

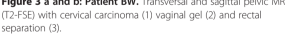
### Results

Hydrogel application was uneventful in all patients and no late toxicity was reported. On axial and sagittal T2weighted MRI hydrogel was clearly visible as a hyperintense region between anterior rectal wall and posterior vaginal wall (Figures 1, 2, 3, 4 and 5). Extend of separation ranged from 7 to 26 mm in wideness (median 10 mm). The length of the separation varied between 18 and 38 mm (median 32 mm), (Table 1). In all patients displacement was seen only in the posterior vaginal fornix and/or deepest part of uterine cervix depending on the extension of the cul-de-sac in correlation to the posterior wall of uterus. In patients with bulky tumor or with deep vaginal extend of peritoneal cavity tumour



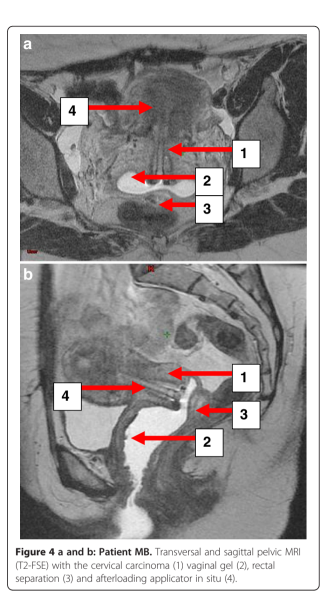
**Figure 2** Instillation of the gel into the Douglas Space during general anaesthesia, # application set with the two components of the gel.





was seen mainly cranial from the rectovaginal space and therefore above the hydrogeI application (Figures 3, 4, 6, 7). Only in the extra-peritoneal part of the cervix a good separation could be achieved between the rectum and cervix (Figure 5).

After completion of primary RCTX patients underwent gynaecologic examinations every 3 months. Additionally post radiation diagnostic curettage was done in order to exclude local recurrence at least 3 month after completion of therapy. In 4/5 patients no vital tumor cells were histologically found and they are free of disease 12 months after therapy. One patient had residual tumour in the cervix and synchronous pulmonary metastases.



# Discussion

Most data concerning rectal separation using hydrogel were published on prostate cancer patients. First cadaveric study has shown a decrease of rectal volume receiving 70 Gy significantly from 23% baseline value to 15%, 4% and 0% with 5 mm, 10 mm and 15 mm separation, respectively [20]. These data could be confirmed by other authors in clinical use during EBRT demonstrating a dose reduction to the anterior rectal wall by 50% [21]. This was associated with decreased risk for acute rectal toxicity compared with historical control [22] and improved quality of life [24]. Even for BT Prada et al. [22] could demonstrate that measured rectal dose for HDR boost was significantly lower by about 2 Gy in prostate cancer patients.

To our best knowledge, the present study is the first on cervical cancer patients with hydrogel instillation.

Similarly to prostate cancer patients, the idea behind was to separate the rectum from the tumour infiltrated cervix uteri and thus reducing radiation dose to the anterior rectal wall during both external beam and brachytherapy planning. Alternative hydrogel distension could allow dose escalation without increasing risk for rectal toxicity.

sagittal orientation with cervical carcinoma (1) vaginal gel (2) and

rectal separation (3).

There are two anatomical characteristis in female pelvis. In contrast to male anatomy with narrow and tense

Table 1	l ength	and	wideness	of	separation	(cm)
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Length of separation (cm)	Wideness of separation (cm)
3.15	1.30
1.79	1.02
3.49	2.61
2.61	0.71
3.77	1.04
	1.79 3.49 2.61

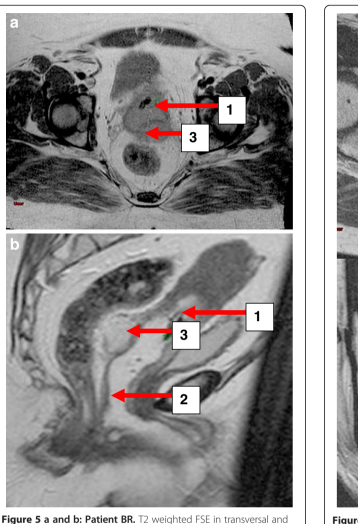
space between rectum and prostate (rectal fascia, Denonville fascia) rectovaginal septum is larger and contains more loose tissue. More volume of hydrogel would be necessary to develop entire rectovaginal space. However, median separation in our pilot study was 10.5 mm,

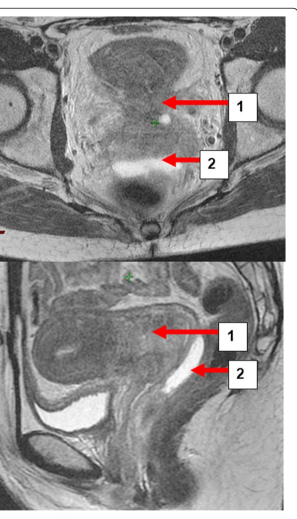
tains more loose tissue. More volume of hydrogel would be necessary to develop entire rectovaginal space. However, median separation in our pilot study was 10.5 mm, which is in the range of the reported values from prostate cancer patients [20]. The length of separation has not been reported by other authors.

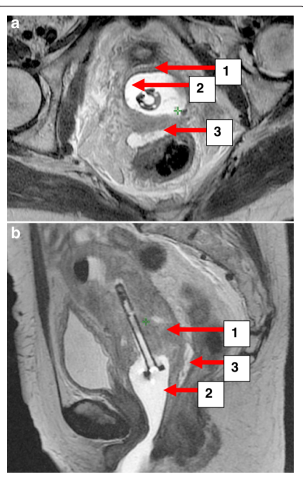
Second anatomical difference is the extent of the culde-sac in correlation to uterine/vaginal posterior wall which varies individually. In contrast to all studies on prostate that showed useful rectal separation, for cervical cancer we could only demonstrate a good separation for the extraperitoneal part of the cervix and upper vagina but no distension in the peritoneal part. This fact limits the value of the method for brachytherapy applications as well as EBRT. If by use of more hydrogel (approximately 100-150c) peritoneal cavity could be elevated and

Figure 6 a and b: Patient LC. Transversal and sagittal pelvic MRI (T2-FSE) with cervical carcinoma (1) and rectal separation (2).









**Figure 7 a and b. Patient MB.** Transversal and sagittal pelvic MRI (T2-FSE) with the cervical carcinoma (1) vaginal gel (2) and rectal separation (3).

therefore rectum as well as small bowel could be more separated, remains speculative. Just as by other authors there was no toxicity from the instillation procedure in our study [20-24].

# Conclusion

Because of different anatomy in the female pelvis injection of 20 cc hydrogel does not seem to be a useful tool for rectal separation and thus for rectal dose sparing in patients with cervical cancer.

#### Competing interests

The author declare that they have no competing interests.

#### Authors' contributions

SM: Data acquisition and Paper Writing; Revision and corresponding author, VB: Idea and Concept, FW: Patient Management, EB: Patient Management, CK: Surgery and Gel Instillation, Manuscript Revision, BG: MRI scans, Diagnostics, Manuscript Revision. All authors read and approved the final manuscript.

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#### References

- Green JA, Kirwan JM, Tierney JF, Symonds P, Fresco L, Collingwood M, Williams CJ: Survival and recurrence after concomitant chemotherapy and radiotherapy for cancer of the uterine cervix: a systematic review and meta-analysis. *Lancet* 2001, 358:781–86.
- Beck T, Sukumvanich P, Rubatt J, Beriwal S, Zorn K, Richard S, Edwards R, Krivak T: Impact of the National Cancer Institutes clinical announcement on cervical cancer survival. *Gynecol Oncol* 2010, 116(Suppl):140.
- Beadle BM, Jhingran A, Yom SS, Ramirez PT, Eifel PJ: Patterns of regional recurrence after definitive radiotherapy for cervical cancer. Int J Radiat Oncol Biol Phys 2010, 76:1396–403.
- Kirwan JM, Symonds P, Green JA, Tierney J, Collingwood M, Williams CJ: A systematic review of acute and late toxiciy of conomitant chemoradiation for cerical cancer. *Radiother Oncol* 2003, 68:217–26.
- Parker K, Gallop-Evans E, Hanna L, Adams M: Five years' experience treating locally advanced cervical cancer with concurrent chemoradiotherapy and high-dose-rate brachytherapy: results from a single institution. Int J Radiat Oncol Biol Phys 2009, 1(74):140–6.
- Chung YL, Jian JJ, Cheng SH, Hsieh CI, Tan TD, Chang HJ, Hung CF, Horng CF, Soong T, Tsou MH: Extended-field radiotherapy and high-dose-rate brachytherapy with concurrent and adjuvant cisplatin-based chemotherapy for locally advanced cervical cancer: a phase I/II study. *Gynecol Oncol* 2005, 97:126–35.
- Chen SW, Liang JA, Hung YC, Yeh LS, Chang WC, Lin WC, Yang SN, Lin FJ: Concurrent weekly cisplatin plus external beam radiotherapy and highdose rate brachytherapy for advanced cervical cancer: a control cohort comparison with radiation alone on treatment outcome and complications. Int J Radiat Oncol Biol Phys 2006, 66:1370–7.
- Small W Jr, Winter K, Levenback C, Iyer R, Gaffney D, Asbell S, Erickson B, Jhingran A, Greven K: Extended-field irradiation and intracavitary brachytherapy combined with cisplatin chemotherapy for cervical cancer with positive para-aortic or high common iliac lymph nodes: results of ARM 1 of RTOG 0116. Int J Radiat Oncol Biol Phys 2007, 68:1081–7.
- Chen MF, Tseng CJ, Tseng CC, Kuo YC, Yu CY, Chen WC: Clinical outcome in posthysterectomy cervical cancer patients treated with concurrent Cisplatin and intensity-modulated pelvic radiotherapy: comparison with conventional radiotherapy. Int J Radiat Oncol Biol Phys 2007, 67:1438–44.
- Heron DE, Gerszten K, Selvaraj RN, et al: Conventional 3D conformal versus intensity-modulated radiotherapy for the adjuvant treatment of gynecologic malignancies: a comparative dosimetric study of dose– volume histograms. Gynecol Oncol 2003, 91:39–45.
- Gerszten K, Colonello K, Heron D, et al: Feasibility of concurrent cisplatin and extended field radiation therapy (EFRT) using intensity-modulated radiotherapy (IMRT) for carcinoma of the cervix. *Gynecol Oncol* 2006, 102:182–8.
- 12. Portelance L, Chao KS, Grigsby PW, *et al*: Intensity- modulated radiation therapy (IMRT) reduces small bowel, rectum and bladder doses in patients with cervical cancer receiving pelvic and paraaortic radiation. *Int J Radiat Biol Oncol Phys* 2001, **51**:261–6.
- D'Souza WD, Ahamad AA, Iyer RB, Salehpour MR, Jhingran A, Eifel PJ: Feasibility of dose escalation using intensity-modulated radiotherapy in posthysterectomic cervical carcinoma. *Int J Radiat Oncol Biol Phys* 2005, 61:1062–70.
- Marnitz S, et al: Helical Tomotherapy with Simultaneous Integrated Boost After Laparoscopic Staging in Patients with Cervical Cancer: Analysis of Feasibility and Early Toxicity. Int J Radiat Biol Phys 2011, [Epub ahead of print].

- D'Souza WD, Ahamad AA, Iyer RB, Salehpour MR, Jhingran A, Eifel PJ: Feasibility of dose escalation using intensity-modulated radiotherapy in posthysterectomic cervical carcinoma. Int J Radiation Oncol Biol Phys 2005, 61:1062–70.
- Adli M, Mayr NA, Kaiser HS, et al: Does prone positioning reduce small bowel dose in pelvic radiation with intensity-modulated radiotherapy for gynecologic cancer? Int J Radiat Ocol Biol Phys 2003, 57:230–8.
- Huh SJ, Kang MK, Han Y: Small bowel displacement system-assisted intensity-modulated radiotherapy for cervical cancer. *Gynecol Oncol* 2004, 93:400–6.
- Chen SW, Liang JA, Hung YC, Yeh LS, Chang WC, Lin WC, Yang SN, Lin FJ: Late toxicities in concurrent chemoradiotherapy using high-dose-rate intracavitary brachytherapy plus weekly cisplatin for locally advanced cervical cancer: a historical cohort comparison against two previous different treatment schemes. Eur J Gynaecol Oncol 2010, 31:504–9.
- Georg P, Lang S, Dimopoulos JC, Dörr W, Sturdza AE, Berger D, Georg D, Kirisits C, Pötter R: Dose-volume histogram parameters and late side effects in magnetic resonance image-guided adaptive cervical cancer brachytherapy. Int J Radiat Oncol Biol Phys 2011, 79:356–62.
- Susil RC, McNutt TR, DeWeese TL, Song D: Effects of prostate-rectum separation on rectal dose from external beam radiotherapy. Int J Radiat Oncol Biol Phys 2010, 76:1251–8.
- Noyes WR, Hosford CC, Schultz SE: Human Collagen Injections to Reduce Rectal Dose During Radiotherapy. Int J Radiat Oncol Biol Phys 2011, Apr 22. [Epub ahead of print].
- 22. Prada PJ, Fernández J, Martinez AA, de la Rúa A, Gonzalez JM, Fernandez JM, Juan G: Transperineal injection of hyaluronic acid in anterior perirectal fat to decrease rectal toxicity from radiation delivered with intensity modulated brachytherapy or EBRT for prostate cancer patients. *Int J Radiat Oncol Biol Phys* 2007, **69**:95–102.
- Wilder RB, Barme GA, Gilbert RF, Holevas RE, Kobashi LI, Reed RR, Solomon RS, Walter NL, Chittenden L, Mesa AV, Agustin J, Lizarde J, Macedo J, Ravera J, Tokita KM: Cross-linked hyaluronan gel reduces the acute rectal toxicity of radiotherapy for prostate cancer. Int J Radiat Oncol Biol Phys 2010, 77:824–30.
- Wilder RB, Barme GA, Gilbert RF, Holevas RE, Kobashi LI, Reed RR, Solomon RS, Walter NL, Chittenden L, Mesa AV, Agustin JK, Lizarde J, Macedo JC, Ravera J, Tokita KM: Cross-linked hyaluronan gel improves the quality of life of prostate cancer patients undergoing radiotherapy. *Brachytherapy* 2011, 10:44–50.
- Marnitz S, Köhler C, Burova E, Wlodarczyk W, Jahn U, Grün A, Budach V, Stromberger C: Helical Tomotherapy with Simultaneous Integrated Boost After Laparoscopic Staging in Patients with Cervical Cancer: Analysis of Feasibility and Early Toxicity. Int J Radiat Oncol Biol Phys 2011, May 18. [Epub ahead of print].

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