Brachytherapy quality assurance in the PORTEC-4a trial for molecular-integrated risk profile guided adjuvant treatment of endometrial cancer

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Objective: The PORTEC-4a trial investigates molecular-integrated risk profile guided adjuvant treatment for endometrial cancer. The quality assurance programme included a dummy run for vaginal brachytherapy prior to site activation, and annual quality assurance to verify protocol adherence. Aims of this study were to evaluate vaginal brachytherapy quality and protocol adherence.

Methods: For the dummy run, institutes were invited to create a brachytherapy plan on a provided CT-scan with the applicator in situ. For annual quality assurance, institutes provided data of one randomly selected brachytherapy case. A brachytherapy panel reviewed and scored the brachytherapy plans according to a checklist.

Results: At the dummy run, 15 out of 21 (71.4%) institutes needed adjustments of delineation or planning. After adjustments, the mean dose at the vaginal apex (protocol: 100%; 7 Gy) decreased from 100.7% to 99.9% and range and standard deviation (SD) narrowed from 83.6–135.1 to 96.4–101.4 and 8.8 to 1.1, respectively. At annual quality assurance, 22 out of 27 (81.5%) cases had no or minor and 5 out of 27 (18.5%) major deviations. Most deviations were related to delineation, mean dose at the vaginal apex (99.9%, 74.7–114.2, SD 7.6) or reference volume length.

Conclusions: Most feedback during the brachytherapy quality assurance procedure of the PORTEC-4a trial was related to delineation, dose at the vaginal apex and the reference volume length. Annual quality assurance is essential to promote protocol compliance, ensuring high quality vaginal brachytherapy in all participating institutes.

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The primary treatment for women with endometrial cancer (EC) is abdominal or laparoscopic hysterectomy with bilateral salpingo-oophorectomy, followed by adjuvant radiotherapy depending on clinicopathological risk factors. Currently, four risk groups of EC have been defined: low, intermediate, high-intermediate (HIR) and high-risk [1].

For women with HIR EC the standard adjuvant treatment is vaginal brachytherapy (VBT), which is based on previous randomised trials. VBT was shown to be equally effective compared to external beam radiotherapy (EBRT) in local control and survival, with a markedly lower toxicity profile [2–6]. However, there is still considerable overtreatment, as approximately 7–10 women with...
HIR EC need to be treated with adjuvant VBT to prevent one recurrence [7]. Better selection of patients at risk of recurrence may play an important role in reducing overtreatment.

The Cancer Genome Atlas Group (TCGA) has discovered four specific molecular subgroups of EC, with each subgroup having a distinct prognosis [8]. Using surrogate markers, these molecular subgroups have been validated in independent EC cohorts and have been shown promising in guiding decisions on adjuvant treatment [9–12]. The role of molecular factors in decision making on adjuvant treatment of HIR EC is currently being investigated in the ongoing international randomised PORTEC-4a trial [13]. In this trial, women with HIR EC are stratified in a favourable, intermediate or unfavourable profile based on molecular and clinicopathologic risk factors and consequently treated with no adjuvant treatment, VBT or EBRT, respectively [14].

In view of the use of VBT in the standard arm and for the women with intermediate profile in the experimental arm, approximately 60% of the PORTEC-4a trial population will receive VBT, a single channel brachytherapy plan using a vaginal cylinder. The VBT planning is based on delineation of the target volume and organs at risk on CT- or MRI-images during at least one fraction. Imaging with CT or MRI with a vaginal cylinder in situ can provide valuable data on dose distribution to the target volume and rectum and bladder that can be used for evaluation of VBT related toxicity. Since institutes had limited experience with delineating on CT- or MRI-scans for single channel VBT, and to ensure uniform high-quality brachytherapy in the PORTEC-4a trial, a dedicated VBT quality assurance (QA) programme, including a dummy run procedure, was implemented in the trial. Especially for radiotherapy trials in general, QA is considered essential as a decrease in therapeutic effectiveness and impaired trial outcomes by protocol deviations have been reported [15, 16]. Furthermore, QA increases trial protocol adherence and treatment uniformity, and therewith ensures optimal treatment in both arms which leads to more reliable trial outcomes [17–23].

The aim of the current study was to investigate protocol adherence by evaluating results of the dummy run procedure and three annual QA rounds in the international PORTEC-4a trial.

Methods

Trial objective

The main objective of the randomised PORTEC-4a trial is to evaluate adjuvant treatment directed by molecular-integrated risk profiles for women with HIR EC, defined as: either (1) FIGO stage IA (with invasion) and grade 3; (2) FIGO stage IB grade 1 or 2 with age ≥60 and/or LVSI; (3) FIGO stage IB grade 3 without LVSI; or (4) FIGO stage II (microscopic) and grade 1. Based on three risk profiles, women in the experimental arm will receive either no further treatment when favourable, adjuvant VBT when intermediate, or EBRT when unfavourable. Women randomised to the standard arm receive adjuvant VBT. Details on patient selection, treatment and trial logistics have been published previously [13, 14].

Trial registration numbers – clinicaltrials.gov (NCT03469674); ISRCTN11659025; NTR8541.

Vaginal brachytherapy in the PORTEC-4a trial

Vaginal brachytherapy should start within 6–8 weeks from the date of surgery. High dose rate (HDR) brachytherapy is given with a vaginal cylinder with one active central channel. Prior to cylinder insertion vaginal examination should take place to verify if the surgical scar has healed sufficiently. Preferably the cylinder with the largest diameter that fits comfortably is used to ensure optimal contact with the vaginal mucosa, resulting in an optimal dose gradient at the surface. After the cylinder placement, correction to a horizontal position is recommended to avoid unnecessary dose to the rectum or bladder [24].

At the first brachytherapy session a CT- or MRI-scan with the applicator in situ is made for delineation of the CTV and organs-at-risk (OARs) and treatment planning. The CTV consists of the vaginal wall and apex of the upper 1/3 of the vagina; for the majority of patients this corresponds to a length of approximately 3.5 cm. The CTV is delineated as a ring structure that surrounds the applicator with a 3 mm margin. OAR include the bladder, rectum, sigmoid and small bowel (loops).

For treatment planning, a library of standard plans per applicator type, diameter and target length are used, with 6 dose reference points, A1 to A6 (Fig. 1). Points A1 and A3 are located at the top of the cylinder at 5 mm from the cylinder surface, with A1 at the central axis and A3 5 mm laterally from A1. Parallel to the central axis at 5 mm from the cylinder surface points A2 and A4 to A6 are placed. A2 is located halfway along the length of the active dwell positions, A4 at the first possible dwell position and point A5 and A6 in between A4 and A2 and caudal of A2, respectively (Fig. 1).

Three fractions of 7 gray (Gy), prescribed to dose point A2, should be delivered within an overall treatment time of 2 weeks. To ensure an adequate dose in the apical vaginal mucosa and compensate for the anisotropy in the longitudinal direction of the 192-Iridium source, the dose in point A1 should be at least 90% and in A3 110% at maximum, with an average dose in A1 and A3 of 100% (7 Gy). A symmetrical loading pattern of the cylinder in the cranio-caudal direction is recommended to facilitate treatment planning, but not mandatory. The reference volume length (RVL) represents the length of the vaginal wall that receives 100% or more and is measured from the top of the 100% isodose line to the point where it enters the cylinder caudally. The RVL should be around 40–45 mm, with a maximum of 50 mm, ensuring sparing of the lower vaginal wall. The mean doses to 90% and 98% of the CTV (D90 and D98) and the maximum dose to 2 cc (D2cc) of the OARs, should be recorded.

Brachytherapy QA-procedure

Dummy run procedure

Before site activation, all participating institutes must have filled in a pre-trial credentialing questionnaire and have performed a dummy run procedure. The questionnaire addresses items such as imaging modality, type of afterloader, cylinder and treatment planning software (TPS) and VBT staff. For the dummy run DICOM-images of a pelvic CT and MR scan with a cylinder in situ are sent to each institute. The local brachytherapy teams are requested to delineate the CTV and OAR conforming to the trial protocol, and to create a brachytherapy plan by using their own TPS. This plan is evaluated by a central QA-panel consisting of two radiation oncologists and one medical physicist specialised in brachytherapy (R.A.N.; C.L.C.; E.A.), a radiation oncologist in training (B.G.W.) and an advanced practitioner brachytherapy (M. S.L.). In case of protocol deviations feedback is sent and the dummy run procedure is repeated when necessary. Upon successful completion of the dummy run procedure, institutes can be activated for the trial.

Annual quality assurance

Annual QA consists of evaluation of a VBT plan of one randomly selected PORTEC-4a case number that has received VBT in the trial in the specific centre. The local team is asked to provide the anonymised CT- or MRI-scan that was used for VBT planning, the DICOM RT-structures, planning and dose distribution, including dose to the A-points, OARs and CTV. Alongside the DICOM-data,
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Updated credentialing questionnaires are requested to objectify changes in VBT components or staff. All requested data, images and plans were evaluated by the QA-panel.

Analysis

According to a QA-checklist all plans of both the dummy run and annual QA were scored on delineation, treatment planning and dose distribution. Annual QA was additionally scored on applicator positioning. Results for each of the items were categorised as fully compliant, partly compliant, in case of a minor protocol deviation, or not compliant, in case of a major deviation. When one or multiple items were scored as partly or not compliant at the dummy run, a revised VBT plan was requested and evaluated. In case of major deviations at annual QA, a teleconference was held for additional explanation and discussion of the feedback, and the next new case number of that particular institute was requested for an extra QA.

On all received data of both the dummy run and the three annual QA procedures descriptive analyses were performed for evaluation of protocol compliance, by comparing the first and final dummy run plan and the annual QA, for the following dose parameters: mean percentage dose, with 100% being 7 Gy, the dose range and standard deviations of all A-points, D98 and D90 of the CTV and the D2cc of the OARs.

To estimate the influence of inter-observer delineation variation on the dose parameters, all delineated structures of the accepted plan of dummy run were projected on the dummy run CT-scan with the LUMC applicator reconstruction and LUMC dose plan. This resulted in the same dose distributions for each case, but varying delineations of the CTV and OARs. For this sub-analysis the mean dose, dose range and standard deviation were recorded. The two institutes with MRI were not included in this analysis.

Results

Between June 1st, 2016 and March 30th, 2020, 327 patients have been included in the PORTEC-4a trial in 19 institutes in 5 countries. Currently, 21 institutes have successfully completed the dummy run. For the dummy run, 19 institutes used CT for brachytherapy planning and two MRI. Three different types of treatment planning systems and three different HDR afterloaders are used (Table 1). Institutes reported the use of several types of single channel vaginal applicators, varying from standard applicators produced by Elekta or Varian, to dedicated applicators, produced in their own institution.

In total, 21 institutes successfully completed the dummy run procedure and participate in the PORTEC-4a trial. Six out of 21 (28.6%) VBT plans were accepted after the first run, 15 (71.4%) needed to resubmit for minor or major adjustments. Common aspects for revisions were: CTV or OAR delineation (Fig. 2A and B), dose planning (Fig. 2C–E) and applicator reconstruction. After adjusting delineation of the CTV and/or dose planning of the VBT plans, the mean dose in the dose prescription point (A2) decreased from 101.5% to 100.5% of the prescribed dose, with 7 Gy being 100%, and the range and standard deviation (SD) of the mean narrowed from 100.0–109.7% to 99.5–105.4% and 2.9 to 1.3, respectively. For the dose at the vaginal apex (mean dose in A1 + A3) the mean decreased from 100.7% to 99.9%, the range from 83.6–135.1% to 96.4–101.4%, and the SD from 8.8 to 1.1 (Table 2). In Table 3 the effect of inter-observer delineation variation on the dose parameters is displayed.

Three annual QA rounds have been performed between September 2017 and February 2020, for which 7, 13 and 7 VBT plans were evaluated in the first, second and in the first part of the third round, respectively. Of 27 requested VBT plans, 22 (81.5%) were accepted with no or minor feedback, while for five (18.5%) plans a teleconference was held for discussion of the feedback, and a new VBT plan of a subsequent case was requested. Most common items for feedback were: CTV delineation (n = 16; CTV length longer than 4.0 cm, or not delineated as a ring structure), or with a margin of more than 3 mm), average dose in points A1 and A3 other than 100% (n = 13; 5 partly (100% ± 3%) and 8 not compliant (100% ± >3%), see Fig. 2C and D), and RVL of more than 50 mm (n = 19, see Fig. 2E and F). Other feedback items addressed applicator positioning (n = 8), suboptimal contact with the vaginal mucosa (n = 5; air or contrast surrounding the applicator), and delineation of the OAR (n = 10, see Table 4).

| Table 1 | Brachytherapy characteristics at dummy run. |
|---------|----------------------------------------|
| Number of institutes |
| Dummy run accepted |
| First plan | 6 |
| Final plan  | 15 |
| Imaging modality |
| CT | 19 |
| MRI | 2 |
| Brachytherapy planning system |
| Oncentra | 14 |
| Flexiplan | 3 |
| Brachyvision  | 4 |
| Type of afterloader |
| Flexitron | 10 |
| Microselectron | 6 |
| Gammamed | 5 |
The treated volumes in the annual QA are displayed in Table 2. The mean dose in the dose prescription point (A2) was 100.4% (range 99.0–108.7%, SD 1.7), and at the vaginal apex (mean A1 + A3) 98.0% (range 74.7–114.2%, SD 7.6). The mean RVL was 53.8 mm, ranging from 44.3 to 70.0 mm. Mean D90 and D98, respectively, of the CTV were 7.9–8.0 Gy and 7.2–7.3 Gy, respectively, in both the dummy run and the annual QA rounds. The mean D2cc of the rectum ranged from 6.0 to 6.1 Gy in both dummy run and QA, and the mean D2cc of the bladder, sigmoid and small bowel varied from 5.2 to 5.9 Gy, 2.9 to 3.7 Gy and 2.8 to 5.5 Gy, respectively (Table 2).

Several changes have been observed in the QA-questionnaires: two institutes changed to a different cylinder applicator, two to another type of afterloader, and three institutes changed their TPS. In five institutes there was a change of brachytherapy staff; two medical physicists and three radiation oncologists were replaced.

**Discussion**

Analysis of the dummy run procedure for vaginal brachytherapy in the PORTEC-4a trial showed that 71.4% of the initially submitted VBT plans needed adjustments to fulfill trial protocol requirements. With the revised VBT plans, an increase in protocol adherence and a decrease in inter-observer delineation and/or dose planning variability were observed, which resulted in more uniform VBT plans. Evaluation of the annual QA of randomly selected VBT plans per centre showed that 18.5% had major protocol deviations, suggesting that a successful dummy run procedure does not rule out major protocol deviations during the trial.
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Table 2
Dose parameters at dummy run procedure and annual QA.

| Dose parameters | Dummy run first plan* (N = 21) | Dummy run final plan (N = 21) | Annual QA (N = 27) |
|-----------------|-------------------------------|-------------------------------|------------------|
| A2 (Aim 100%)   |                               |                               |                  |
| Mean dose (SD)  | 101.5 (2.9)                   | 100.5 (1.3)                   | 100.4 (1.7)      |
| Range           | 100.0–109.7                   | 99.5–105.4                    | 99.0–108.7       |
| A1 (Aim 90–95%) |                               |                               |                  |
| Mean dose (SD)  | 93.1 (8.8)                    | 92.0 (1.7)                    | 90.2 (7.0)       |
| Range           | 75.8–126.8                    | 89.3–94.8                     | 67.8–102.9       |
| A3 (Aim 105–110%) |                               |                               |                  |
| Mean dose (SD)  | 108.2 (9.0)                   | 107.8 (1.8)                   | 105.7 (8.4)      |
| Range           | 91.3–143.4                    | 102.5–110.0                   | 81.7–125.5       |
| A4              |                               |                               |                  |
| Mean dose (SD)  | 83.2 (7.1)                    | 84.9 (6.9)                    | 87.1 (6.2)       |
| Range           | 73.8–106.0                    | 76.6–99.0                     | 76.7–99.1        |
| A5 (Aim 95–100%)|                               |                               |                  |
| Mean dose (SD)  | 96.4 (3.2)                    | 96.8 (3.7)                    | 98.5 (2.9)       |
| Range           | 91.0–103.8                    | 93.0–110.2                    | 94.0–105.1       |
| A6 (Aim 95–100%)|                               |                               |                  |
| Mean dose (SD)  | 97.3 (3.2)                    | 96.9 (3.1)                    | 98.7 (3.5)       |
| Range           | 88.9–102.1                    | 88.9–102.1                    | 87.6–104.6       |
| D90*            |                               |                               |                  |
| Mean (SD)       | 7.9 (0.8)                     | 8.0 (0.9)                     | 8.0 (0.9)        |
| Range           | 6.3–9.4                       | 6.3–9.4                       | 5.6–9.7          |
| D98             |                               |                               |                  |
| Mean (SD)       | 7.2 (0.9)                     | 7.3 (0.9)                     | 7.3 (1.2)        |
| Range           | 5.1–8.6                       | 5.8–8.6                       | 4.0–9.2          |
| Bladder D2cc    |                               |                               |                  |
| Mean (SD)       | 5.2 (0.5)                     | 5.3 (0.5)                     | 5.9 (0.8)        |
| Range           | 4.3–6.0                       | 4.3–6.0                       | 4.8–7.7          |
| Rectum D2cc     |                               |                               |                  |
| Mean (SD)       | 6.1 (0.5)                     | 6.1 (0.5)                     | 6.0 (0.6)        |
| Range           | 4.7–7.1                       | 5.0–6.9                       | 4.0–7.2          |
| Sigmoid D2cc    |                               |                               |                  |
| Mean (SD)       | 3.7 (1.2)                     | 3.6 (1.2)                     | 2.9 (1.2)        |
| Range           | 2.1–6.9                       | 1.4–6.4                       | 0.5–5.1          |
| Small bowel D2cc|                               |                               |                  |
| Mean (SD)       | 5.5 (1.6)                     | 5.3 (1.8)                     | 2.8 (1.8)        |
| Range           | 1.1–7.8                       | 0.9–7.3                       | 0.8–6.9          |

* Institutes for which the first dummy run plan was accepted have been listed in both columns (N = 6).
** Mean percentage dose, with 100% being 7 Gy.
*** Dose in Gy.

In this quality assurance study, most common reasons for feedback were delineation of the CTV and OAR, the average dose at the vaginal apex (dose points A1 + A3) and the reference volume length (RVL). Dose points A1 and A3 represent the vaginal vault area which is essential for the target volume. These dose points are aimed to obtain a uniform and reproducible dose distribution at 5 mm from the apex, even with use of different types of cylinders, sources and treatment planning systems in a randomised multicentre trial. The dose at the apex is essential, not only because approximately over 75% of all recurrences occur at the vaginal apex, but also because a higher dose in point A3 could lead to increased toxicity due to the adjacent bowel loops [25–27]. The RVL directly displays the actual length of the vaginal wall receiving 100% of the dose. The mean RVL in this study was 53.8 mm, while following the trial protocol, the RVL should range between 40 and 50 mm. In case of an increased RVL, a longer segment of the vagina receives significant dose. This observation led to a general feedback to make all participating institutes aware of this and re-emphasise the importance of the trial planning aims.

Minor feedback items addressed the applicator placement and applicator diameter. When the applicator was placed ventrally or dorsally this could lead to higher doses to the bladder or rectum [24]. In 5 out of 27 reviewed cases the diameter of the vaginal applicator seemed relatively small and air gaps or contrast surrounded the applicator, directly affecting the dose distribution. A previous study showed an average dose reduction to the vaginal mucosa of 27% when air gaps were present and stressed that air gaps of more than 2 mm can lead to a decrease in dose to the vaginal mucosa, which in turn may result in an increased risk of local recurrence. Institutes were provided feedback to ensure that an attempt is made to reposition the applicator or to use a larger diameter applicator for more optimal contact to the vaginal mucosa. However, the presence of air gaps has not been related to clinical outcome, as a wide range of dose and fractionation schedules for VBT has been proven effective [28–31].

Data of dose parameters showed improvements in the dose range between the first and the final plan of the dummy run.

Table 3
Variation in dose parameters resulting from inter-observer differences in delineation at the dummy run.

| Dose parameter | Mean*(SD) | Range |
|----------------|----------|-------|
| CTV D90       | 8.1 (0.5) | 7.4–9.2 |
| CTV D98       | 7.3 (0.6) | 6.3–8.4 |
| Rectum D2cc   | 5.9 (0.5) | 4.7–6.8 |
| Bladder D2cc  | 5.0 (0.6) | 4.4–5.6 |
| Sigmoid D2cc  | 3.5 (0.9) | 2.1–6.1 |
| Small bowel D2cc | 5.6 (0.8) | 4.1–6.8 |

* Dose in Gy.

Table 4
Evaluation of the annual QA.

| Items                              | Fully compliant | Partly compliant* | Not compliant* |
|------------------------------------|-----------------|------------------|----------------|
| Applicator positioning             |                 |                  |                |
| Position and angle of cylinder     | 19              | 2                | 6              |
| Contact of cylinder to vaginal     | 22              | 1                | 4              |
| mucosa                             |                 |                  |                |
| Delineation                        |                 |                  |                |
| CTV delineation                    | 11              | 11               | 5              |
| OAR delineation                    | 17              | 8                | 2              |
| Treatment planning-Reconstruction  |                 |                  |                |
| Position of A points               | 24              | 1                | 2              |
| Prescribed dose in point A2         | 22              | 3                | 2              |
| Symmetry of loading pattern        | 18              | 0                | 9              |
| Evaluation of dose distribution    |                 |                  |                |
| Average dose in A1 + A3 = 100%     | 14              | 5                | 8              |
| Dose in point A1 ≥ 90% and/or A3 ≥ 110% | 17          | 6                | 4              |
| Reference length/width              | 8               | 15               | 4              |
| CTV D90/D98                        | 19              | 1                | 7              |
| OAR D2cm3                          | 24              | 3                | 0              |

* Scored according to the detailed description in the trial protocol.
procedure for the essential dose points A1, A2 and A3, indicating the increased protocol adherence. However, at annual QA, one or more years after the initial dummy run, an increased variability in dose distribution and in dose to points A1, A2, A3 and A6 was observed, also at institutes with a large case load. Possible explanations for this could be institutional changes in type of applicator, afterloader, TPS or VBT staff, that were recorded in the questionnaires; adherence to a local VBT protocol; unfamiliarity with CTV delineation for single channel VBT or unfamiliarity with the trial protocol due to infrequent inclusion. This indicates that continuous QA is essential to ensure protocol adherence in the years after the initial dummy run.

The range and standard deviation of dose parameters D90 and D98 of the CTV and D2cc of the OAR remained similar in the first and final plan of the dummy run, even after adjustments of the delineation and/or VBT planning. This could be explained by the impact of delineation variations on these parameters. Additional analysis showed that when eliminating treatment planning variation, by projecting delineations of all institutes on one standard VBT plan, similar standard deviations and ranges were found for CTV D90/98 and D2cc of the OAR in the accepted plans. This means that this remaining variability in dose parameters is caused by inter-observer delineation variations and this should be taken into account when interpreting dose parameter data. Contouring of organs at risk on MRI scans would have been more precise than on CT-scans, but only a minority of centres have MRI available for standard cylinder-based brachytherapy.

Using a uniform protocol for VBT ensures high quality VBT and is essential for increasing reliability of dose parameters that can be used for evaluation of VBT related toxicity. A continuous QA-programme in a multi-institutional radiotherapy trial can increase treatment and delineation uniformity and which has been shown to impact on trial outcomes [15,32]. A review on QA for radiotherapy in randomised trials showed that major protocol deviations were observed in 11.0–48.0% of all cases, and were reported to be associated with impaired overall survival and local control and potentially increased treatment related toxicity [15]. This has also been reported by several other investigators, emphasising that the design of the QA-procedure needs to be tailored to specific trial techniques and outcomes [20,22,23,32–34].

To our knowledge, this is the first study on dedicated QA for single channel VBT with delineation on CT- or MRI-scans for endometrial cancer. Our findings confirm that a dummy run and QA-procedure in multi-institutional radiotherapy trials creates awareness of the trial protocol and principles and guidelines of the specific treatment, improves protocol adherence and quality of the treatment. Even after successful initial dummy run procedures, annual QA showed major protocol deviations in 18.5% of reviewed cases, suggesting that continuous annual QA is essential to promote protocol adherence, ensuring uniform high-quality vaginal brachytherapy a multi-institutional trial.

Conflicts of interest

The authors do not have conflicts of interest to disclose.

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References

[1] Colombo N., et al., ESMO-ESGO-ESTRO Consensus Conference on Endometrial Cancer: diagnosis, treatment and follow-up. Ann Oncol, 2016, 27: 16–41.
[2] Sorbe B et al. Intravaginal brachytherapy in FIGO stage I low-risk endometrial cancer: a controlled randomized study. Int J Gynecol Cancer 2009;19:873–8.
[3] Nout RA et al. Vaginal brachytherapy versus pelvic external beam radiotherapy for patients with endometrial cancer of high-intermediate risk (PORTEC-2): an open-label, non-inferiority, randomised trial. Lancet 2010;375:816–23.
[4] Sorbe B et al. External pelvic and vaginal irradiation versus vaginal irradiation alone as postoperative therapy in medium-risk endometrial carcinoma—a prospective randomized study. Int J Radiat Oncol Biol Phys 2012:82:1249–55.
[5] de Boer SM et al. Long-term impact of endometrial cancer diagnosis and treatment on health-related quality of life and cancer survivorship: results from the randomized PORTEC-2 trial. Int J Radiat Oncol Biol Phys 2015:93:797–809.
[6] Wortman BG et al. Ten-year results of the PORTEC-2 trial for high-intermediate risk endometrial carcinoma: improving patient selection for adjuvant therapy. Br J Cancer 2018;119:1067–74.
[7] Thomas GM. A role for adjuvant radiation in clinically early carcinoma of the endometrium?. Int J Gynecol Cancer 2010;20(Suppl 2):564–6.
[8] Kandoth C et al. Integrated genomic characterization of endometrial carcinoma. Nature 2013;497:67–73.
[9] Talhouk A et al. A clinically applicable molecular-based classification for endometrial cancers. Br J Cancer 2015;113:299–310.
[10] Stelloo E et al. Improved risk assessment by integrating molecular and clinicopathological factors in early-stage endometrial cancer—combined analysis of the PORTEC cohorts. Clin Cancer Res 2016;22:4215–24.
[11] Talhouk A et al. Confirmation of ProMisE: A simple, genetics-based clinical classifier for endometrial cancer. Cancer 2017;123:802–13.
[12] Kommoss S et al. Final validation of the ProMiSe molecular classifier for endometrial carcinoma in a large population-based case series. Ann Oncol 2018;29:1180–8.
[13] Creutzberg CL, PORTEC-4a: molecular profile-based versus standard adjuvant radiotherapy in endometrial cancer (PORTEC-4a). https://clinicaltrials.gov/ct2/show/NCT03469674, 2016. Accessed March 27, 2020.
[14] Wortman BG et al. Molecular-integrated risk profile to determine adjuvant radiotherapy in endometrial cancer: evaluation of the pilot phase of the PORTEC-4a trial. Gynecol Oncol 2018.
[15] Weber DC et al. QA makes a clinical trial stronger: evidence-based medicine in radiation therapy. Radiother Oncol 2012;105:4–8.
[16] Ohri N et al. Radiotherapy protocol deviations and clinical outcomes: a meta-analysis of cooperative group clinical trials. J Natl Cancer Inst 2013;105:387–93.
[17] Ibbott GS et al. Challenges in credentialing institutions and participants in advanced technology multi-institutional clinical trials. Int J Radiat Oncol Biol Phys 2008;71(1 Suppl):S136–41.
[18] Cormack RA. Quality assurance issues for computed tomography-, ultrasound-, and magnetic resonance imaging-guided brachytherapy. Int J Radiat Oncol Biol Phys 2008;71(1 Suppl):S136–41.
[19] Bekelman JE et al. Redesigning radiotherapy quality assurance: opportunities to develop an efficient, evidence-based system to support clinical trials—report of the National Cancer Institute Work Group on Radiotherapy Quality Assurance. Int J Radiat Oncol Biol Phys 2012;83:782–90.
[20] Fairchild A et al. Do results of the EORTC dummy run predict quality of radiotherapy delivered within multicentre clinical trials?. Eur J Cancer 2012;48:3232–9.
[21] Ibbott GS, Haworth A, Followill DS. Quality assurance for clinical trials. Front Oncol 2013;3:311.
[22] Fairchild A et al. Quality assurance for the EORTC Z2071–26071 study: dummy run prospective analysis. Radiat Oncol 2014;9:248.
[23] Kirstis C et al. Quality assurance in MR image guided adaptive brachytherapy for cervical cancer: Final results of the EMBRACE study dummy run. Radiat Oncol 2015;11:548–54.
[24] Hoskin PJ, Bowens P, Summers A. The influence of applicator angle on dosimetry in vaginal vault brachytherapy. Br J Radiol 2002;75:234–7.
[25] Creutzberg CL et al. Surgery and postoperative radiotherapy versus surgery alone for patients with stage-I endometrial carcinoma: multicentre randomised trial. PORTEC Study Group. Post Operative Radiation Therapy in Endometrial Carcinoma. Lancet 2000;355:1404–13.
[26] Keys HM et al. A phase III trial of surgery with or without adjunctive external pelvic radiotherapy in intermediate risk endometrial adenocarcinoma: a Gynecologic Oncology Group study. Gynecol Oncol 2004;92:744–51.
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[27] Hoskin P, et al. The GEC-ESTRO handbook of brachytherapy: endometrial cancer. 2016.
[28] Cameron AL, Cornes P, Al-Booz H. Brachytherapy in endometrial cancer: quantification of air gaps around a vaginal cylinder. Brachytherapy 2008;7:355–8.
[29] Richardson S, Palaniswamy G, Grigsby PW.Dosimetric effects of air pockets around high-dose rate brachytherapy vaginal cylinders. Int J Radiat Oncol Biol Phys 2010;76:276–9.
[30] Humphrey P, Cornes P, Al-Booz H. Vaginal vault brachytherapy in endometrial cancer: verifying target coverage with image-guided applicator placement. Br J Radiol 2013;86:20120428.
[31] Pearcey RG, Petereit DG. Post-operative high dose rate brachytherapy in patients with low to intermediate risk endometrial cancer. Radiother Oncol 2000;56:17–22.
[32] Fairchild A et al. Does quality of radiation therapy predict outcomes of multicenter cooperative group trials? A literature review. Int J Radiat Oncol Biol Phys 2013;87:246–60.
[33] Poortmans PMP et al. The potential impact of treatment variations on the results of radiotherapy of the internal mammary lymph node chain: a quality-assurance report on the dummy run of EORTC phase III randomized trial 22922/10925 in stage I-III breast cancer. Int J Radiat Oncol Biol Phys 2001;49:1399–408.
[34] Ibbott GS et al. Dose specification and quality assurance of RTOG protocol 95–17: a cooperative group study of 192Ir breast implants as sole therapy. Int J Radiat Oncol Biol Phys 2007;69:1572–8.