Objective

The aim of the present study was to determine if volumetric modulated arc therapy (VMAT) provided superior dose distribution than intensity-modulated radiotherapy (step-and-shoot; ssIMRT) based on target volume coverage and organs at risk (OARs) doses in postoperative radiotherapy for pancreatic cancer patients.

Methods

New 4-dimensional computed tomography plans for 10 pancreatic cancer patients were created. The ssIMRT plans had 6 coplanar fields (330-0-30-60-90°) and VMAT plans were generated with 2 268-92° arcs.

Results

VMAT plans revealed better overall sparing of right kidney (volume receiving 15% of prescribed dose [V15]: 28.3% vs 46.9%, p=0.012; V20: 16.1% vs 27.6%, p=0.007; V25: 8.6% vs 15.2%, p=0.005; mean dose 1549 centigray [cGy] vs 1987 cGy, p=0.005). VMAT delivered similar isodose distribution (planning target volume [PTV] mean dose: 5164 vs 5183 cGy, PTV max: 5526 cGy vs 5505 cGy; p=0.541) with significantly fewer monitor units (MU) (MU: 468 vs 527; p=0.032) in comparison with ssIMRT. VMAT was also found to be superior for V30 intestinal dose, but mean dose was similar (1963 cGy vs 2032 cGy; p=0.05).

Conclusion

VMAT provided more effective protection for bilateral kidneys and small intestine with better OAR doses, as well as for liver, with reduced high-dose volumes in this cohort. This could be investigated as more tolerable concurrent radiochemotherapy treatment with better OAR preservation.

Keywords: Intensity-modulated radiation therapy; pancreatic cancer; volumetric modulated arc therapy.

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therapy in the United States, while “chemotherapy alone” is the current standard in Europe.[1] One of the important reasons for this divergence especially in the postoperative setting, is the location of pancreas as it is located in the middle of multiple critical structures like kidneys, small bowel, liver and spinal cord which leads more radiotherapy toxicity.[2]

Intensity-modulated radiation therapy (IMRT) is an advanced mode of high-precision radiotherapy that delivers radiation doses precisely to the three-dimensional shape of the tumor by modulating the intensity of the radiation beam in multiple small volumes while minimizing the dose to surrounding normal critical structures.[3] In the postoperative or curative setting, compared to 3-D Conformal planning, IMRT has encouraging gastrointestinal toxicity results in treating pancreas cancer without compromising local control.[4–8] In the dosimetric evaluation by Landry et al., compared to 3-D-CRT, using IMRT reduced the amount of small bowel receiving more than 50 Gy form 31% to 19.2%.[5] First clinical result from Wayne state University reported 7% incidence of grade 3 gastrointestinal toxicity and one year actuarial survival rate of 69% for postoperative or locally advanced pancreas cancer patients treated with 54–55 Gy IMRT.[6] Systematic review of clinical results comparing IMRT to 3DCRT have also shown the reduction of grade 2 and 3 toxicities without compromising of local control.[9]

The volumetric modulated arc therapy (VMAT) technique is a recent form of IMRT with using one or two-arc gantry rotation by simultaneously modulating the MLC position and the dose rates. In the recent reports, VMAT has been shown to be superior for a variety of cancer types such as head and neck, prostate cancer and the dosimetric and clinic publications seeking for new cancer sites to use VMAT were increasing.[10,11] As the motion is an important problem in radiotherapy planning of pancreas cancer, VMAT can decrease the risk of intrafractional organ motion by shortening treatment time.[12] In the dosimetric studies, comparing VMAT to Helical tomotherapy, IMRT and 3D-CRT, reduced organ at risk doses such as kidney and liver have a promising effect on reducing toxicity due to radiotherapy.[13,14]

We aimed to define whether VMAT provides a superior dose distribution in comparison to Intensity modulated radiotherapy (Step and shoot: ssIMRT) based on 4D-CT target volume coverage and organs at risk (OAR) doses in adjuvant postoperative radiotherapy for pancreas adenocarcinoma patients.

Materials and Methods

Patients

We have re-planned the planning 4D CT (Respiratory data sets are “binned” by phase: 0–100% at 10% interval) scans of our ten consecutive stage IIB pancreas cancer patients in 2011. All patients were treated with postoperative concurrent radiochemotherapy of 50.4 Gy in 28 fractions (ssIMRT, 1.8 Gy/fraction/day) with Capecitabine 2500 mg/m². Whipple surgery for pancreatic head tumors was performed.

Simulation and target contouring

All the patients were simulated in the supine position on a customized vacuum bed, with using T-bar, Wing-board, and knee-foot stopper. 4-D and contrast free-breathing axial CT scans with 3 mm slice thickness were obtained by AcQSim CT simulation of Philips Brilliance Big Bore CT. Respiratory correlated imaging was generated for planning which was performed by Pinnacle radiation therapy planning system (9.0, Philips Medical Systems Inc. Cleveland, OH) which uses Collapse Cone [cc] convolution algorithm. Also for every patient a second scan with intravenous contrast were obtained and fused for contouring purposes.

Planning

Clinical target volume (CTV) was delineated according to departmental guidelines and RTOG pancreas cancer web based CT contouring atlas,[8] as porta hepatitis, superior mesenteric artery, celiac, paraaortic and postoperative bed including peripancreatic nodal area based on preoperative CT fusion. An internal CTV as ITV, showing the position of CTV in all 4D-CT scans were generated by one physician and the second physician reviewed all the contours. Planning target volume (PTV) was integrated tumor volume (ITV=internal CTV contoured on all respiratory data sets) plus 4 mm. A “small-bowel region” was defined which consisted of the abdominal content after subtracting the PTV, all OARs and the vertebral bodies, with the posterior border extending to the dorsum of the lumbar vertebral body, but excluding the retroperitoneal space.[12] Liver was delineated at MinIP phase.

The prescribed dose was 50.4 Gy in 28 fractions and the planning objective was to give at least 95% prescribed dose to PTV and 98% prescribed doses to CTV. Identical objectives were used for IMRT and VMAT plans. The planning objectives were selected as listed; The maximum point dose to spinal cord is less than 45 Gy, volume of kidney receiving more than 20 Gy (V20)
<33%, if one kidney exceeds the above, then spare the other kidney with 20 Gy (V20) <20%, mean liver dose (MLD) <32Gy, V20Gy <66%.

Treated ssIMRT plans were 6 coplanar fields (330-0-30-60-90 degree) with multiple segments. VMAT plans were generated as two 268–92° arcs rotating clockwise and counter clockwise starting from 92° and 268° with 15° collimator angle. Collimator angle was fixed to -15° to minimize the effects of interleaf leakage and tongue-and groove effect. For all the plans, 6 MV Photon beams created by using Varian Linac Triology (Rapid-Arc) 120 leaf millennium multileaf collimator (MLC) with a maximum dose rate 600 MU/min and Grid Size was 0.3x0.3x0.3 cm for both plan calculations. All plans were performed by one physicist (YS).

**Comparison of VMAT and ssIMRT techniques**
The maximum dose (Gy) for spinal cord, V15-V20-V25 for bilateral kidney, V30 and mean dose for intestine, V30-V45 for liver, total treated monitor units (MU), and mean and maximum doses for PTV were compared based on dose volume histograms. VMAT and ssIMRT techniques were compared using two-tailed pair wise Wilcoxon signed-ranked tests, with p<0.05 considered to indicate statistically significant differences.

**Results**
The mean ITV volume in 10 patients was 462.0 cm³ (range: 286.8–793.5 cm³). The mean doses to liver, both kidneys and other critical structures were listed in Table 1.

**Target**
In all the plans PTV was covered at least 95% prescribed dose of 50.4Gy. VMAT delivered similar isodose distributions (PTV mean dose: 5164 vs 5183 cGy, PTV Max 5526 cGy vs 5505 cGy, p=0.541). As an advantage VMAT has been delivered by significantly less MU (MU: 468 vs 527, p=0.032) in comparison to ssIMRT. Representative isodose curves for ssIMRT and VMAT technique were given in axial, sagittal and coronal CT slices at Figures 1–3.

Kidneys: Right kidney was spared better as V15-28.3% vs 46.9%, p=0.012; V20-16.1% vs 27.6%, p=0.007; V25- 8.6% vs 15.2%, p=0.005; mean dose 1594cGy vs 1987 cGy, p=0.005 compared to 6 field ssIMRT. For left kidney, there was decrease in high dose regions (V20 11.6% vs 18.8%; p=0.008, V25 5.7% vs 11.7%; p=0.018). No statistical difference was found in terms of mean doses (1155 cGy vs 1209 cGy, p=0.33) and V15 -low dose area (V15; 23.4% vs 29.8%, p=0.74) for left kidney.

**Table 1** Dosimetric comparisons of IMRT and VMAT planning techniques for postoperative pancreas cancer patients

<table>
<thead>
<tr>
<th>Parameter</th>
<th>VMAT mean dose</th>
<th>VMAT range</th>
<th>ssIMRT mean</th>
<th>ssIMRT range</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Left kidney</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>V15 (%)</td>
<td>23.4</td>
<td>4.8–41</td>
<td>29.8</td>
<td>7.3–56.</td>
<td>0.74</td>
</tr>
<tr>
<td>V20 (%)</td>
<td>11.6</td>
<td>1.5–30</td>
<td>18.8</td>
<td>3.2–32</td>
<td>0.018</td>
</tr>
<tr>
<td>V25 (%)</td>
<td>5.7</td>
<td>0.3–14</td>
<td>11.7</td>
<td>1.3–24</td>
<td>0.01</td>
</tr>
<tr>
<td>Mean dose (cGy)</td>
<td>1155</td>
<td>695–1617</td>
<td>1209</td>
<td>808–1771</td>
<td>0.33</td>
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<tr>
<td>Right kidney</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>V15 (%)</td>
<td>28.3</td>
<td>9–49</td>
<td>46.9</td>
<td>28–62</td>
<td>0.012</td>
</tr>
<tr>
<td>V20 (%)</td>
<td>16.1</td>
<td>5–30</td>
<td>27.6</td>
<td>19–39</td>
<td>0.007</td>
</tr>
<tr>
<td>V25 (%)</td>
<td>8.6</td>
<td>1–18</td>
<td>15.2</td>
<td>9–27</td>
<td>0.001</td>
</tr>
<tr>
<td>Mean dose (cGy)</td>
<td>1549</td>
<td>811–3832</td>
<td>1987</td>
<td>1300–4853</td>
<td>0.002</td>
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<tr>
<td>Liver</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>V30 (%)</td>
<td>18.3</td>
<td>10–30</td>
<td>20.1</td>
<td>12–30</td>
<td>0.012</td>
</tr>
<tr>
<td>V45 (%)</td>
<td>7.9</td>
<td>5–14</td>
<td>9.5</td>
<td>6–15</td>
<td>0.006</td>
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<tr>
<td>Mean dose (cGy)</td>
<td>1507</td>
<td>1094–2320</td>
<td>1550</td>
<td>1097–2113</td>
<td>0.168</td>
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<td>Small bowel</td>
<td></td>
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<td></td>
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<tr>
<td>V30 (%)</td>
<td>28.1</td>
<td>19–48</td>
<td>32.9</td>
<td>22–59</td>
<td>0.008</td>
</tr>
<tr>
<td>Mean dose (cGy)</td>
<td>1963</td>
<td>1009–2963</td>
<td>2032</td>
<td>1017–3278</td>
<td>0.05</td>
</tr>
<tr>
<td>Spinal cord maximum dose (cGy)</td>
<td>3697</td>
<td>3199–4100</td>
<td>3792</td>
<td>3456–4233</td>
<td>0.24</td>
</tr>
<tr>
<td>PTV mean dose (cGy)</td>
<td>5183</td>
<td>5097–5260</td>
<td>5164</td>
<td>5096–5195</td>
<td>0.06</td>
</tr>
<tr>
<td>PTV maximum dose (cGy)</td>
<td>5505</td>
<td>5370–5971</td>
<td>5526</td>
<td>5325–5973</td>
<td>0.54</td>
</tr>
<tr>
<td>Monitor units</td>
<td>468</td>
<td>319–624</td>
<td>527</td>
<td>420–583</td>
<td>0.032</td>
</tr>
</tbody>
</table>

ssIMRT: Step and shoot intensity modulated radiotherapy; VMAT: Volumetric modulated arc radiotherapy.
comparable spinal cord doses (cord maximum dose 3792 cGy vs 3697 cGy, p=0.24).

Discussion

Radiotherapy for pancreas cancer is a challenging area due to its location and motion factor because of respiratory and other organ movements. In attempts to dose escalation in the adjuvant setting, Allen et al., reported that maximum tolerated dose was 39 Gy using 3DCRT

Small Bowel; VMAT was also found to be superior for intestinal doses (V30: 28.1% vs 32.9%, p=0.008) but both planning technique provides similar mean doses of small bowel.

Liver; the mean doses were found to be similar in each plan, but high dose volumes such as V30 and V45 were decreased by VMAT (VMAT vs ssIMRT: mean dose- 1507 cGy vs 1550, p=0.168, V30- 18.3% vs 20.1%, p=0.012, V45- 7.9% vs 9.5%, p=0.006).

Spinal cord; Both plans provided acceptable and comparable spinal cord doses (cord maximum dose 3792 cGy vs 3697 cGy, p=0.24).

Fig. 1. Representative isodose curves for step and shoot intensity modulated radiotherapy (ssIMRT) and volumetric arc therapy (VMAT) technique. 54 Gy curve-red, 45 Gy curve-orange 35 Gy curve-yellow and 25 Gy curve-blue were shown in axial slices.

Fig. 2. Representative isodose curves for step and shoot intensity modulated radiotherapy (ssIMRT) and volumetric arc therapy (VMAT) technique. 54 Gy curve-red, 45 Gy curve-orange 35 Gy curve-yellow and 25 Gy curve-blue were shown in coronal slices.
with concurrent gemcitabine, and escalation to 42 Gy resulted increased gastrointestinal toxicity.[15] Also the gastrointestinal tumor study group (GITSG) trial that lead the treatment approach used today, delivered 40 Gy split course radiotherapy with a 2 weeks break.[16] IMRT has been shown to improve dosimetric parameters and decrease normal tissue doses; as a result it has been accepted as a routine approach. The first important problem of radiotherapy is the high rated of gastrointestinal acute side effects during concurrent radiotherapy. While older series using 3DCRT has almost 58% non-hematological grade 3 or more acute toxicity rate, newer series using IMRT has a rate ranging from 7% to 16%. [6,16–18] The second concern was interfraction and intrafraction movement in pancreas cancer, using VMAT which is a novel form of IMRT which offers fast and homogeneous dose delivery and 4DCT simulation, could provide more effective dose distribution and achieve to maintain the dose delivery as it was planned. In our study, based on 4DCT scans, VMAT allowed dose reduction in right kidney, liver, small bowel and high dose volume of left kidney compared to ssIMRT. To our knowledge, this is the first study that compares VMAT to IMRT based on 4D-CT scans of a pure patient population whom underwent Whipple operation.

One of the important acute side effect was caused by the dose that small bowel received during therapy. It is shown that even large volume of low dose areas (5–15 Gy) also can cause toxicity.[19] This is a concern with using novel technologies, this is not supported by clinical studies evaluating acute GI side effects.[18] Landry et al., delivered 45 Gy for gross tumor volume and microscopic disease, and in comparison with 3DCRT, IMRT decreased the dose that one third of the small bowel receiving from 38.5 Gy vs. 30.2 Gy. Also IMRT reduced the median volume of small bowel receiving more than 50 Gy.[5] Milano et al., analyzed 25 patients who was treated for pancreas and bile duct cancers with concurrent chemoradiotherapy using IMRT technique. Of the patients, 80% has grade 2 or less acute upper GI toxicity.

Memorial Sloan Kettering Cancer center has evaluated 205 patients that were treated for locally advanced pancreas cancer undergoing IMRT or 3DCRT. The evaluation of Grade 2+ GI toxicity revealed a reduction from 34% to 16% by using IMRT compared to using 3DCRT which shows the clinical benefit of using advanced planning.[4] There results were supported by a systematic review of published clinical studies consisting 45 patients comparing IMRT and 3DCRT where grade 3 toxicities were dropped from 10.6% to 5% (p=0.017). In our study VMAT was also found to be superior to IMRT for high dose region (V30) but no significant difference for the mean small bowel doses with using 4DCT planning.

Brown et al., compared IMRT, integrated boost IMRT and 3DCRT for locally advanced pancreas cancer and revealed that left kidney was spared better in 3DCRT due to the use of multiple fields in IMRT, in contrast to the right kidney doses which was better with IMRT.[7] In other comparison of IMRT to 3DCRT, V20 right kidney doses were lowered from 27.7% to 16%, but this not observed for left kidney (5.7% vs 11.1%, p=0.1).[20] The superiority of IMRT over 3DCRT has been also shown dosimetrically by Bahl et al., where V45 for bowel bag was 212.3±159.0 cc (mean volume±standard deviation).
expiration or end-inspiration was found to have limited benefits in sparing normal organs.[20] Sangalli et al., studied the impact of 4DCT planning for unresectable pancreatic cancer patients. The results revealed that by using 4DCT, target volumes reduced by 37% compared to standard target delineation.[25] In our protocol, 4D CT simulation was performed as a standard clinic routine for planning of pancreas cancer patients.

Conclusions
In the dosimetric comparison of ssIMRT and VMAT techniques based on 4D CT scans of postoperative pancreas cancer patients, both plans reach dosimetric organ goals, but VMAT provided superior dose distribution in terms of organs at risk such as bilateral kidneys and small intestine, as well as liver with reduced high dose volumes in this cohort without compromising CTV coverage. This study is limited by a relatively small number of patient's 4DCT scans but these results could be promising for more tolerable concurrent radiochemotherapy treatments and increasingly usage of VMAT for pancreas cancer as routine like the other sites such as head and neck.

Disclosure Statement
The authors declare no conflicts of interest.

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