An Investigation of Non-Coplanar Volumetric Modulated Radiation Therapy for Locally Advanced Unresectable Pancreatic Cancer Using a Trajectory Optimization Method

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Purpose: This study was conducted in order to develop a trajectory optimization algorithm for non-coplanar volumetric modulated arc therapy (VMAT) and investigate the potential of organs at risk (OARs) sparing in locally advanced pancreatic cancer patients using non-coplanar VMAT.

Methods and Materials: Firstly, a cost map that represents the ray–OAR voxel intersections at each source position was generated using a ray-tracing algorithm. A graph search algorithm was then used to determine the least-cost path from the cost map. Lastly, full arcs or partial arcs were selected based on the least-cost path to generate the non-coplanar VMAT (ncVMAT) trajectories. Clinical coplanar VMAT (coVMAT) plans for 11 patients diagnosed with locally advanced unresectable pancreatic cancer (LAPC) receiving 45 to 70 Gy in 25 fractions were replanned using non-coplanar VMAT trajectories. Both coplanar and non-coplanar plans were normalized to cover 95% of the PTV45 Gy volume with a prescription dose of 45 Gy. The conformity index (CI), homogeneity index (HI), PTV coverage, and dose to the OARs were compared between coVMAT and ncVMAT plans.

Results: With ncVMAT, the mean coverage of PTV50 Gy, PTV54 Gy, PTV60 Gy, and PTV70 Gy increased significantly. The mean conformity index of PTV45 Gy, PTV54 Gy, and PTV70 Gy was also improved in the ncVMAT plans. Compared with coVMAT plans, the ncVMAT plans resulted in significantly lower doses to the spinal cord, bilateral kidneys, stomach, and duodenum. The maximum dose to the spinal cord decreased by 6.11%. The mean dose to the left and right kidneys decreased by an average of 5.52% and 11.71%, respectively. The Dmax, Dmean, and D15% of the stomach were reduced by an average of 7.45%, 15.82%, and 16.79%, separately. The D15% and Dmean of the duodenum decreased 6.38% and 5.64%, respectively.
INTRODUCTION

According to the American Cancer Society, pancreatic cancer was the seventh leading cause of cancer-related death worldwide for both genders in 2018 (1). The only curative therapy for patients diagnosed with pancreatic cancer is surgical resection. Nevertheless, only 15%–20% of these patients presented with resectable tumors (2). The standard care for patients with unresectable pancreatic cancer is chemotherapy and/or chemoradiotherapy (3). However, conventionally fractionated radiotherapy which delivers 40 to 60 Gy in 1.8–2.0 Gy per fraction showed minimal to no local tumor control benefit for patients with locally advanced pancreatic cancer (LAPC) (4). With the development of image-guided radiotherapy and immobilization techniques, stereotactic body radiotherapy (SBRT) which precisely delivers a high dose per fraction has become a promising option for the treatment of patient with LAPC. A study that investigated the outcomes of patients treated with SBRT using the National Cancer Data Base (NCDB) showed that the median overall survival (OS) (13.9 months) and the 2-year OS rate (21.7% vs. 11.6 months) and the 2-year OS rate (21.7% vs. 16.5%) were significantly higher with SBRT versus conventionally fractionated radiation therapy (5). The reason why SBRT resulted in a better survival rate is that SBRT allows to deliver a higher biological effective dose (BED) to patients and higher BED is associated with better local control. On the other hand, early studies implementing SBRT were associated with high early and/or late gastrointestinal side effects (6, 7). Therefore, the sparing of radiosensitive gastrointestinal organs such as the stomach, small intestines, colon, and duodenum near the pancreas becomes the main difficulty of dose escalation in LAPC patients (4).

A lot of studies have been done to investigate the application of non-coplanar radiotherapy in OAR dose sparing for different sites. Evidence has shown that non-coplanar plans were superior in normal tissue sparing compared with the conventional coplanar technique (8–12). Smyth et al. evaluated non-coplanar VMAT for OAR sparing in primary brain tumor radiotherapy (13). They found that compared with coplanar VMAT, non-coplanar VMAT significantly reduced doses to the contralateral lobe, optic nerve, hippocampus, and temporal lobe for patients with primary brain tumors. Yu et al. reported the first clinical implementation of 4π radiation therapy in recurrent high-grade glioma patients (14). Twenty beam orientations were used for optimization. The results showed that compared with coplanar VMAT, 4π IMRT plans resulted in equal or significantly lower OAR doses. A study shows that in head and neck cases, non-coplanar IMRT can significantly reduce dose to hippocampi and whole brain which helps preserve neurocognitive function (15). Another study of non-coplanar VMAT for brain metastases shows that non-coplanar VMAT generated more rapid dose falloff and higher conformity compared with co-planar VMAT (8). Uto et al. compared coplanar VMAT and non-coplanar VMAT for hippocampus protection in craniopharyngiomas cases and found that non-coplanar VMAT significantly reduced the dose to bilateral hippocampus (16). The two non-coplanar arcs were set at couch angles of 45° and 315°. Similarly, Cheung et al. applied non-coplanar VMAT to primary brain tumor using non-coplanar arcs with couch rotated to 45° or 315° (17). An alternative approach to non-coplanar IMRT or non-coplanar VMAT is the VMAT+ technique proposed by Sharfo et al. which combined full coplanar arcs with few non-coplanar beams with optimized beam angles (18). The advantage of the VMAT+ technique is that it has similar plan quality to fully non-coplanar IMRT but much shorter treatment delivery time.

Non-coplanar techniques have also been implemented in the management of pancreatic cancer. Chang et al. found that non-coplanar IMRT was able to significantly decrease bilateral kidney dose for unresectable pancreatic cancer compared with coplanar IMRT (19). The non-coplanar IMRT beam angles were manually selected in beam’s eye view (BEV) to spare the kidneys. Osborne et al. also found that the non-coplanar plans show overall benefits over coplanar plans (20). Two lateral wedged oblique fields and two wedged non-coplanar oblique fields were used. Burghelea et al. investigated non-coplanar trajectories for LAPC cases using dynamic wave arc on the VERO gimballed linac (21). However, the beam directions were selected by a human planner. Previous studies on pancreatic cancer were based on the IMRT technique or the beam angles were selected manually due to the complexity of beam orientation optimization (22). In this study, we present an implementation of non-coplanar VMAT technique which utilized cost map generation and trajectory optimization algorithm for OAR dose sparing in LAPC patients.

MATERIALS AND METHODS

Patient Selection

This retrospective study was approved by the institutional review board and informed consent was waived. Eleven patients with locally advanced pancreatic cancer treated with conventional coplanar VMAT plans were selected in our study. Patients were simulated using Philips Brilliance Big-Bore helical CT with 3 mm slice thickness. Plans were designed and optimized using VMAT techniques with simultaneous integrated boost (SIB) in the Eclipse treatment planning system (Varian Medical System, Inc.).
Palo Alto, CA, USA). PTV prescription dose includes 45 Gy in 1.8 Gy fractions, 50 Gy in 2 Gy fractions, 54 Gy in 2.14 Gy fractions, 60 Gy in 2.4 Gy fractions, and 70 Gy in 2.8 Gy fractions. The average planning target volume (PTV) size was 435.31 cm$^3$ for PTV$_{45Gy}$, 215.32 cm$^3$ for PTV$_{50Gy}$, 45.50 cm$^3$ for PTV$_{54Gy}$, 43.53 cm$^3$ for PTV$_{60Gy}$ and 20.37 cm$^3$ for PTV$_{70Gy}$.

**Data Input**

For each patient, the Digital Imaging and Communications in Medicine (DICOM) structure set which contains isocenter position, contours of PTV, and OARs were exported from Eclipse treatment planning system into MATLAB (The MathWorks, Natick, MA, USA) to generate a 3D patient phantom for ray-tracing simulation.

**Ray Tracing and Cost Map Generation**

The gantry rotated from 0° to 360° with control points spaced every 2°, and the couch rotated from 270° to 90° with control points spaced every 2°. For each source position determined by the couch and gantry angle, ray tracing was performed using a MATLAB algorithm (23). The couch and gantry angle combinations which may cause collision were eliminated from ray tracing. The collision regions were determined by measuring the permitted gantry rotation angle for each couch angle with a patient phantom placed on the couch.

For each source position, ray tracing was performed, and a list of coordinates in three dimensions of the intersected voxels was produced. Then, the number of intersected voxels in different types of OARs was calculated. The OARs evaluated in this study were the small intestines, duodenum, colon, liver, bilateral kidneys, stomach, and spinal cord.

For each source position determined by couch angle $c$ and gantry angle $g$, the cost of each node $C_{cg}$ is given by the sum of the relative volume of each OAR intersected during ray tracing:

$$C_{cg} = \sum_{v \in V} \frac{n_v}{N_v}$$

where $v$ is the OAR of interest, $V$ is the set of all OARs evaluated, $n_v$ is the number of organ $v$ voxels intersected by the ray, and $N_v$ is the total number of voxels of organ $v$. After the cost of each permitted source position was calculated, the cost data were reformatted into a matrix and displayed as a cost map as shown in Figure 1A.

**Trajectory Selection**

After the cost map was generated, the trajectory optimization was simplified as a traveling salesman problem. A salesman wants to travel between several cities to sell goods. The distance to each city is different and not every city is connected. The problem is to find the shortest route between a certain number of destinations that must be visited. One constraint of the traveling salesman problem is that each city can be visited only once. Similar to the salesman problem, we want to find the least-cost trajectory through various nodes. The cost of each node is similar to the distance to each city in the traveling salesman problem. Dijkstra’s algorithm was used to find the minimal cost trajectory through the cost map. Dijkstra’s algorithm aims to find the shortest paths from a starting node to all other nodes in a weighted graph. This algorithm was created by the Dutch computer scientist Dr. Edsger W. Dijkstra. In this study, we only need to find the shortest path from the source node to a target node. The source node was at gantry angle of 0 and couch angle of 0. The target node was at gantry angle of 360 and couch angle of 0. The reason of doing this is that every gantry angle from 0° to 360° will be visited. There are four inputs when implementing Dijkstra’s algorithm. The first input is the cost of moving from node $i$ to node $j$ which is the sum of the cost of node $i$ and node $j$. The second input is an adjacency matrix which indicates which nodes are adjacent to which other nodes in the cost map:

$$A(i,j) = \begin{cases} 1, & \text{if } g_i - g_j = 2^\circ \text{ or } |c_i - c_j| = 2^\circ \\ 0, & \text{else} \end{cases}$$

where $g_i$ and $g_j$ are the gantry angle of node $i$ and node $j$, and $c_i$ and $c_j$ are the couch angle of node $i$ and node $j$. The last two inputs are the coordinates of the source node and target node. The outputs are the minimal cost path connecting the starting node and ending node and the cost value for the path. Figure 1B shows the least-cost path generated by Dijkstra’s algorithm. Due to restrictions of the Eclipse treatment planning system, the couch and gantry cannot rotate simultaneously for VMAT plans. Therefore, the optimized trajectory was divided into several subarcs with fixed couch positions as shown in Figure 1C. Subarcs with an arc length less than 30° cannot be optimized in the treatment planning system and, therefore, were excluded from plan optimization.

**Treatment Planning**

The original clinical coVMAT plans consisted of two coplanar arcs with control points spaced every 2°. Plans were designed and optimized using the Eclipse planning system. Both the coplanar and non-coplanar plans were normalized to cover 95% of PTV$_{45Gy}$ volume with a prescription dose of 45 Gy. Coplanar and non-coplanar plans were produced for a 6MV Varian Trilogy linear accelerator with a Millennium 120 leaf MLC. The minimal MLC width was 5 mm. The dose calculation algorithm was the anisotropic analytical algorithm (AAA). The calculation resolution was 2.5 mm.

**Plan Evaluation**

Multiple dose metrics were evaluated for coVMAT and ncVMAT plans. For the targets, homogeneity index (HI), conformity index (CI), and mean coverage (V100%) were compared.

The HI was defined as:

$$HI = \frac{D_{5\%}}{D_{95\%}} \times 100,$$

where $D_{5\%}$ and $D_{95\%}$ are the doses to 5% and 95% of the PTV volume.
CI was recommended by Paddicks’s formula:

\[
CI = \frac{V(PI \cap T)}{V(T)} \times \frac{V(PI \cap T)}{V(PI)},
\]

where \(V(PI \cap T)\) is the volume of the intersection of the prescription isodoses and the target, \(V(T)\) is the volume of the target, and \(V(PI)\) is the volume of the prescription isodose. For OARs, the maximum point dose (\(D_{\text{max}}\)), dose to 15% of OAR volume (\(D_{15\%}\)), dose to 2 cm\(^3\) of OAR volume (\(D_{2\, cc}\)), the portion of volume receiving 20 Gy (\(V_{20\, Gy}\)), and 40 Gy (\(V_{40\, Gy}\)) were evaluated for the small intestine, colon, and stomach. The mean dose (\(D_{\text{mean}}\)) was evaluated for bilateral kidneys. \(D_{\text{mean}}\) was evaluated for the liver. The \(D_{\text{max}}\) was also evaluated for the spinal cord. \(D_{\text{max}}\) and \(D_{15\%}\) were evaluated for the duodenum. Those criteria were selected according to RTOG 0848 and practice in our institution (24). Table 1 shows the normal tissue dose constraints for conventionally fractionated radiotherapy from RTOG 0848. Monitor unit (MU) was calculated for each plan. Paired t-test was used to perform statistical analysis, and a significance level of \(P \leq 0.05\) (two-tailed) was used.

RESULTS

The dosimetric parameters of the PTVs are shown in Table 2. Both coVMAT and ncVMAT plans were normalized to over 95% of the PTV\(_{45\, Gy}\) volume with the prescription dose of 45 Gy. Therefore, the mean coverage of PTV\(_{45\, Gy}\) was both 95% for coVMAT and ncVMAT plans. Compared with coVMAT plans, the mean coverage of PTV\(_{50\, Gy}\), PTV\(_{54\, Gy}\), PTV\(_{60\, Gy}\), and PTV\(_{70\, Gy}\) of ncVMAT plans were significantly increased. The mean conformity index of PTV\(_{45\, Gy}\), PTV\(_{54\, Gy}\), and PTV\(_{70\, Gy}\) was also increased, while the mean conformity index of PTV\(_{50\, Gy}\) and PTV\(_{60\, Gy}\) decreased slightly. There was no significant difference for homogeneity indexes between the coVMAT and ncVMAT plans.

The dosimetric parameters of the OARs are shown in Table 3. Compared with coVMAT plans, the ncVMAT plans resulted in a lower dose to the liver. The mean dose to the liver was reduced by an average of 8.8%. Compared with coVMAT plans, ncVMAT plans resulted in lower doses to the spinal cord. The max dose to the spinal cord was reduced by an average of 6.11%. The ncVMAT plans also resulted in lower doses to bilateral kidneys. The mean dose to the left kidney decreased by an average of 5.52%. The mean dose to the right kidney decreased by an average of 11.71%. For the stomach, a significant improvement of the \(D_{\text{max}}\), \(D_{\text{mean}}\), and \(D_{15\%}\) was observed. For the small intestines, ncVMAT resulted in lower \(D_{\text{max}}\), \(D_{\text{mean}}\), \(D_{2\, cc}\), and \(V_{20\, Gy}\), while \(V_{40\, Gy}\) increased slightly. There was no significant difference between coVMAT and ncVMAT for small intestine \(D_{15\%}\). The \(D_{\text{mean}}\) and \(V_{20\, Gy}\) of the colon decreased, while the colon \(V_{40\, Gy}\) increased. Colon \(D_{\text{max}}\), \(D_{2\, cc}\), and \(D_{15\%}\) were similar for coVMAT and ncVMAT. Compared with coVMAT plans, ncVMAT plans also resulted in lower \(D_{15\%}\) and \(D_{\text{mean}}\) to the duodenum.

Figures 2, 3 show the dose distribution comparison of coVMAT and ncVMAT for one typical patient and the dose-volume histogram of PTV and selected OARs. As Figure 2 shows, the pancreas is centrally located and surrounded by many radiation-sensitive organs such as the stomach, bilateral kidneys, duodenum, colon, and small intestine. From the dose distribution, we can see that the ncVMAT plan has a better sparing of the liver, small intestine, and colon due to the additional freedom of couch rotation. The dose-volume histogram comparison also shows that ncVMAT resulted in a

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**TABLE 1 | Normal tissue dose constraints.**

| Structure               | RTOG 0848                        |
|-------------------------|----------------------------------|
| Spinal cord             | Max dose <4,500 cGy              |
|                         | Mean dose ≤2,500 cGy             |
| Liver                   | Max dose ≤5,400 cGy              |
| Small bowel             | \(D_{15\%}\) ≤4,500 cGy         |
| Stomach                 | Max dose ≤5,400 cGy              |
|                         | \(D_{15\%}\) ≤5,000 cGy         |
|                         | Mean dose ≤1,800 cGy             |
| Bilateral kidneys       |                                 |
|                         |                                 |

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**FIGURE 1 | (A) Cost map generated through ray-tracing simulation. (B) The least cost trajectory generated by Dijkstra’s algorithm displayed on cost map with collision region excluded. (C) The subarcs with fixed couch positions (displayed as red lines) adopted as final non-coplanar trajectories.**

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significant dose reduction to selected OARs with comparable target coverage to the coVMAT plan. In such a case, the improved OAR sparing can be utilized to increase the amount of dose that can be delivered to the tumor limited by normal tissue toxicity, thus improving the local control.

The average MUs for coVMAT and ncVMAT plans were 585 ± 117 and 540 ± 80 MU. The average number of subarcs for each non-coplanar plan is 10. The minimum and maximum numbers of subarcs are 8 and 12.

DISCUSSION

In the radiotherapy of pancreatic cancer, a hypofractionated regimen that delivers much higher BED to the tumor shows better local control compared with the conventionally fractionated regimen. However, the implementation of hypofractionated radiotherapy in pancreatic cancer is usually associated with excess normal tissue toxicity which limited the amount of radiation dose that can be delivered to the tumor. Thus, OAR sparing is of great significance in order to improve tumor control while reducing normal tissue toxicity. In this study, non-coplanar VMAT was investigated for its ability of OAR dose sparing in locally advanced pancreatic cancer patients by using a trajectory optimization approach. Due to the complexity of non-coplanar VMAT trajectory optimization, beam orientations were selected manually in previous studies. The quality of non-coplanar plan using manually selected beam direction largely relies on the experience of the treatment planner. This study investigates the organ at risk sparing in locally advanced pancreatic cancer patients using the non-coplanar VMAT technique. Compared with previous studies which manually select the beam orientation, the non-coplanar trajectories in our study were selected automatically using the trajectory optimization algorithm. Compared with automatic optimization methods such as the VMAT+ technique, our method is simple and easy to implement because the fluence map optimization is decoupled from beam orientation optimization. Firstly, a cost map which represents the ray–OAR overlap was produced through ray-tracing simulation for each patient. Then, a shortest path finding algorithm was utilized to find the minimal cost trajectory through the cost map. The result in non-coplanar trajectory was a continuous path. Due to restriction of the planning and delivery system, the linac gantry and couch cannot rotate simultaneously, and the non-coplanar VMAT trajectory was divided into several subarcs with static couch rotation.

For OARs, both the coVMAT and ncVMAT plans met the dose constraints suggested by RTOG 0848. However, ncVMAT

### Table 2 | Dosimetric parameters of targets for coVMAT and ncVMAT plans.

|                | Coplanar | Non-coplanar |
|----------------|----------|--------------|
| PTV45 Gy CI    | 0.87 ± 0.07 | 0.91 ± 0.12  |
| PTV45 Gy HI    | 1.41 ± 0.14 | 1.43 ± 0.15  |
| PTV45 Gy V100%| 95%       | 95%          |
| PTV40 Gy CI    | 0.77 ± 0.10 | 0.68 ± 0.15  |
| PTV40 Gy HI    | 1.37 ± 0.14 | 1.38 ± 0.15  |
| PTV40 Gy V100%| 89.08% ± 6.93% | 93.07% ± 3.99% |
| PTV40 Gy CI    | 0.34 ± 0.28 | 0.42 ± 0.22  |
| PTV40 Gy HI    | 1.02      | 1.02         |
| PTV40 Gy V100%| 44.35% ± 34.35% | 62.95% ± 36.45% |
| PTV60 Gy CI    | 0.84 ± 0.17 | 0.73 ± 0.14  |
| PTV60 Gy HI    | 1.14 ± 0.08 | 1.14 ± 0.08  |
| PTV60 Gy V100%| 96% ± 4.34% | 97% ± 2.86%  |
| PTV60 Gy CI    | 0.73 ± 0.12 | 0.84 ± 0.24  |
| PTV60 Gy HI    | 1.04 ± 0.01 | 1.05 ± 0.01  |
| PTV60 Gy V100%| 89.25% ± 10.68% | 97.38% ± 2.54% |

### Table 3 | Normal tissue dose–volume statistics.

| Dose metrics                  | Coplanar            | Non-coplanar        | P     |
|-------------------------------|---------------------|---------------------|-------|
| Liver mean dose (Gy)          | 11.55 ± 12.51       | 10.66 ± 12.27       | 0.07  |
| Rt kidney mean dose (Gy)      | 9.14 ± 3.96         | 8.16 ± 3.73         | 0.020 |
| Lt kidney mean dose (Gy)      | 12.53 ± 3.37        | 11.74 ± 2.99        | 0.035 |
| Spinal cord max dose (Gy)     | 37.59 ± 6.99        | 35.02 ± 5.95        | 0.067 |
| Small intestine max dose (Gy)| 51.59 ± 3.39        | 50.21 ± 3.58        | 0.029 |
| Small intestine mean dose (Gy)| 12.96 ± 7.22        | 12.08 ± 7.01        | 0.051 |
| Small intestine D2cc (Gy)     | 45.50 ± 7.74        | 43.96 ± 8.82        | 0.035 |
| Small intestine volume >20 Gy (cm³) | 136.14 ± 100.35 | 123.06 ± 86.45 | 0.110 |
| Small intestine volume >40 Gy (cm³) | 31.72 ± 36.92 | 32.95 ± 35.80 | 0.495 |
| Small intestine D15% (Gy)     | 24.44 ± 13.16       | 24.52 ± 13.35       | 0.878 |
| Colon max dose (Gy)           | 46.56 ± 8.74        | 46.09 ± 9.47        | 0.327 |
| Colon mean dose (Gy)          | 12.93 ± 4.19        | 11.89 ± 3.85        | 0.160 |
| Colon D2cc (Gy)               | 40.74 ± 8.79        | 40.82 ± 8.99        | 0.875 |
| Colon volume >20 Gy (cm³)     | 134.22 ± 80.70      | 123.99 ± 84.93      | 0.499 |
| Colon volume >40 Gy (cm³)     | 7.04 ± 6.60         | 11.48 ± 12.04       | 0.108 |
| Colon D15% (Gy)               | 23.08 ± 5.98        | 22.96 ± 6.55        | 0.921 |
| Stomach max dose (Gy)         | 48.34 ± 6.45        | 45.40 ± 9.53        | 0.036 |
| Stomach mean dose (Gy)        | 14.40 ± 8.55        | 12.72 ± 8.48        | 0.004 |
| Stomach D15% (Gy)             | 14.40 ± 8.55        | 12.72 ± 8.48        | 0.004 |
| Duodenum max dose (Gy)        | 45.40 ± 11.71       | 45.55 ± 11.91       | 0.690 |
| Duodenum D15% (Gy)            | 39.31 ± 10.83       | 37.65 ± 12.20       | 0.049 |
| Duodenum mean dose (Gy)       | 30.29 ± 13.75       | 29.18 ± 14.37       | 0.062 |
significantly decreased the dose to the stomach, bilateral kidneys, duodenum, and spinal cord, compared with coVMAT. The stomach $D_{\text{max}}$, $D_{\text{mean}}$, and $D_{15\%}$; the bilateral kidneys $D_{\text{mean}}$; the duodenum $D_{\text{mean}}$ and $D_{15\%}$; and the spinal cord $D_{\text{max}}$ were all decreased with ncVMAT. An overall lower mean dose to the small intestine and colon was also achieved with ncVMAT.

As Figure 4 shows, the non-coplanar VMAT significantly improved the mean coverage of PTV$_{50\ Gy}$, PTV$_{54\ Gy}$, PTV$_{60\ Gy}$ and PTV$_{70\ Gy}$. This could potentially improve tumor control. Improved tumor control also means improved survival. The mean conformity index of PTV$_{45\ Gy}$, PTV$_{54\ Gy}$, and PTV$_{70\ Gy}$ was also improved which indicated that ncVMAT may have a better sparing of normal tissues.

Although the MU for ncVMAT plans was slightly lower than the coVMAT plans, the treatment delivery time of ncVMAT plans can be longer than coVMAT plans due to the extra couch rotation between multiple arcs. However, the treatment time of ncVMAT plans can be reduced by implementing automatic couch rotation.

![Figure 2](image1.jpg)  
**FIGURE 2** | Dose distribution for one typical patient shown in axial, coronal, and sagittal views: (A) coplanar VMAT and (B) non-coplanar VMAT plan.

![Figure 3](image2.jpg)  
**FIGURE 3** | Dose–volume histogram of PTV and selected OARs (solid line—coVMAT, dashed line—ncVMAT).
There are several limitations of this study: 1) OARs were equally weighted when generating the cost map. A customized weighting factor can be applied to each OAR according to their clinically relevant importance to generate better sparing of a specific organ. 2) Trajectory optimization was separated from VMAT plan optimization. Due to the computation complexity, the trajectory optimization was separated from VMAT plan optimization. 3) Beams that had opposite direction were equally weighted during ray tracing. The OAR can be between the source and the PTV or behind the PTV for opposite beam directions. It is less desirable to have the OAR between the source and the PTV. In such a case, the beam orientations which have OAR between the source and the PTV should be heavily penalized. 4) Treatment time and patient comfort were sacrificed by resetting the couch to multiple arc positions.

In summary, compared with conventional coplanar VMAT, non-coplanar VMAT resulted in improved coverage for PTV$_{50}$ Gy, PTV$_{54}$ Gy, PTV$_{60}$ Gy and PTV$_{70}$ Gy. The conformity index of PTV$_{45}$ Gy, PTV$_{54}$ Gy and PTV$_{70}$ Gy was also improved. For OARs, non-coplanar VMAT produced better dose sparing for the stomach, bilateral kidneys, duodenum, and spinal cord. A lower mean dose to the small intestine and colon was also achieved with non-coplanar VMAT with no significant difference for other OARs. In conclusion, non-coplanar VMAT has the potential of increasing the dose delivered to the tumor while reducing normal tissue toxicity, thus improving the local control of locally advanced pancreatic cancer.

**CONCLUSIONS**

A trajectory optimization algorithm was developed for non-coplanar VMAT. Compared with conventional coplanar VMAT, non-coplanar VMAT resulted in improved coverage and conformity to the targets. The sparing of OARs was significantly improved in non-coplanar VMAT compared with coVMAT plans for locally advanced pancreatic cancer.

**DATA AVAILABILITY STATEMENT**

The original contributions presented in the study are included in the article/supplementary material. Further inquiries can be directed to the corresponding author.

**AUTHOR CONTRIBUTIONS**

All the authors have participated sufficiently in the work to take public responsibility for the appropriateness of the material and method and the collection, analysis, and interpretation of the data. All authors contributed to the article and approved the submitted version.

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