Optimizing Patient Selection for Irreversible Electroporation of Locally Advanced Pancreatic Cancer: Analyses of Survival

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Background: Irreversible electroporation (IRE) has emerged as a viable consolidative therapy after induction chemotherapy, in which this combination has improved overall survival of locally advanced pancreatic cancer (LAPC). Optimal timing and patient selection for irreversible electroporation remains a clinically unmet need. The aim of this study was to investigate preoperative factors that may assist in predicting progression-free and overall survival following IRE.

Methods: A multi-institutional, prospectively maintained database was reviewed for patients with LAPC treated with induction chemotherapy followed by open-technique irreversible electroporation from 7/2015-5/2019. RECIST 1.1 criteria were used to assess tumor response and radiological progression. Overall survival (OS) and progression-free survival (PFS) were recorded. Survival analyses were performed using Kaplan Meier and Cox multivariable regression analyses.

Results: 187 LAPC patients (median age 62 years range, 21 – 91, 65% men, 35% women) were treated with IRE. Median PFS was 21.7 months and median OS from diagnosis was 25.5 months. On multivariable analysis, age ≤ 61 (HR 0.41, 95%CI 0.21-0.78, p<0.008) and no prior radiation (HR 0.49, 95%CI 0.26-0.94, p=0.03) were positive predictors of OS after IRE. Age ≤ 61(HR 0.53, 95%CI, 0.28-.99, p=0.046) and FOLFIRINOX followed by gemcitabine/abraxane induction chemotherapy (HR 0.37,95% CI 0.15-0.89, p=0.027) predicted prolonged PFS after IRE. Abnormal CA19-9 values at the time of surgery negatively impacted both OS (HR 2.46, 95%CI 1.28-4.72, p<0.007) and PFS (HR 2.192, 95%CI 1.143-4.201, p=0.018) following IRE.
Conclusions: Age, CA 19-9 response, avoidance of pre-IRE radiation, and FOLFIRINOX plus gemcitabine/abraxane induction chemotherapy are prominent factors to consider when referring or selecting LAPC patients to undergo IRE.

Keywords: locally advanced pancreatic cancer, irreversible electroporation (IRE), overall survival, patient selection, recurrence, progression free survival

INTRODUCTION

The diagnosis of pancreatic ductal adenocarcinoma (PDAC) continues to have a challenging prognosis, but improvements in multi-disciplinary care have raised the overall survival rates to 10% for all stages (1). In 2020, an estimated 47,050 patients will die of this disease representing the third highest cancer causing mortality rate (2). Modern systemic chemotherapy followed by surgical resection has dramatically improved the standard of care, however this is only available to approximately 10-20% of the patients diagnosed each year. Forty percent of patients present with local invasion [stage III - locally advanced pancreatic cancer (LAPC)] and are most often prescribed poorly responsive systemic palliative chemotherapy (3). Multimodality induction chemotherapy with folinic acid, 5-fluorouracil, leucovorin, irinotecan, and oxaliplatin (FOLFIRINOX) has prolonged overall survival (OS) to 12 months, however continued response rates after 4-6 months are poor due to the cumulative toxicity, need for dose delay, dose reduction, or complete termination of this active treatment (4-6). A current clinical unmet need is to develop and offer clinically effective therapies to consolidate the response of systemic chemotherapy in patients with unresectable LAPC after 3-4 months of induction therapy.

Historically, LAPC is deemed unresectable to conventional surgical intervention and is thought of as a continuum of metastatic disease. Yet additional consolidative treatment options for LAPC following induction chemotherapy exist and have been successfully utilized with improved outcomes. Irreversible electroporation (IRE), a non-thermal ablation technology, has begun to gain acceptance within the last decade (7–9) IRE induces cellular apoptosis without disrupting surrounding tissue structural integrity (10). Martin et al. demonstrated IRE is a safe and effective treatment of LAPC with initial improvements in median OS to 25.3 months (11). These results were confirmed with combination of chemotherapy and IRE improving median overall survival to 30.7 months, critically implicating IRE to be included in the multimodal treatment of LAPC (12).

The aim of this study was to evaluate LAPC pre-procedural/preoperative patient predictors of progression-free survival (PFS) and OS following induction chemotherapy to better guide patient selection for IRE utilization as part of a multimodal treatment for LAPC.

METHODS

Participants

An Institutional Review Board (IRB) approved single arm study of patients diagnosed National Comprehensive Cancer Network (NCCN) stage III LAPC of patients treated by IRE between July 2015 and May 2020 was evaluated. This prospective pancreatic cancer registry represents a multi-institutional collection of patients with radiographic stage III LAPC all of whom were treated with IRE (13). Six participating institutions included the University of Louisville, University of South Florida, Augusta University, University of Alabama, and University of California, San Diego. The registry is open to any center worldwide that wishes to participate and collaborate with their data (12). All patients provided written informed consent. A diagnosis of LAPC disease was established by biopsy proven adenocarcinoma of the pancreas with unreconstructable venous involvement or greater than 180° encasement of their superior mesenteric artery (SMA) or celiac artery without evidence of metastatic lesions (12, 14, 15). Patients were also further sub-classified by our recent Stage III classification sub-types (16). Patients were further considered for inclusion in the study if the treating physician at the aforementioned participating institutions believed that ablation of their soft tissue would be feasible in the care of their disease, as has been previously described and outlined (17–19). Staging included triple phase computed tomographic (CT) scan with less than 1.5-mm cuts at the time of diagnosis and repeated 1-2 weeks prior to IRE (11, 20). To aide in post ablation follow up and response, positron emission tomography (PET-CT) scanning was initiated in January of 2019.

Inclusion criteria involved eligible patients underwent induction therapy consisting of chemotherapy and/or external beam radiation therapy following each respective institution’s protocol. Patients underwent restaging evaluation 4 to 6 weeks after induction therapy via repeat triple-phase CT scan and serum tumor markers. Those with evidence of disease progression were excluded. Patients found on restaging to be free of metastatic disease and without primary tumor progression were included and received either IRE in situ or IRE with resection. All patients included were Stage III LAPC based on pre-operative imaging. Patient selection is critical to the safety and efficacy of IRE for LAPC. This has been outlined extensively in previous publications (17, 19, 21).

Key exclusion criteria were patients with implanted cardiac pacemaker or defibrillators unable to be deactivated, non-removable implants with metal parts within 1 cm of the target lesion, a myocardial infarction within 3 months, or unsuitable for general endotracheal anesthesia. All presented data was collected and maintained in a prospective manner. Adverse events were summarized using the National Cancer Institute (NCI) Common Terminology Criteria Adverse Event (CTCAE), version 3.0 and graded via Clavien-Dindo classification (22).

Interventions

Systemic Therapy

FOLFIRINOX based chemotherapy was administered for at least 6 to 8 cycles on a 14-day cycle, commonly using standard dosing
per standard of care and each institutions management (23). Similarly Gemcitabine and abraxane were administered using standard dosing per standard of care and each institutions management on days 1, 8, and 15 every 4 weeks (24). Patients were re-staged after induction chemotherapy via repeat triple-phase CT scan and serum tumor markers and evaluated by a multidisciplinary team. All patients with evidence of disease progression were excluded. Only patients that had received FOLFIRINOX, gemcitabine and abraxane, or single agent gemcitabine as an induction therapy prior to IRE were included in survival analyses.

Irreversible Electroporation

Patients found to be free of metastatic disease and without primary tumor progression on re-staging were included and further received an open surgical in situ IRE based on intra-operative findings and location of the primary tumor as described previously (11, 25). Open Insitu-IRE was performed utilizing AngioDynamics NanoKnife system, as previously described and were performed by surgeons in the operating room (17, 19, 26). All participating institutions utilized the registry protocol for standardization of settings setup and delivery of energy during the IRE procedure as previously reported (12, 17, 19, 27).

Post-Procedure Evaluation and Follow Up

After IRE follow-up imaging via triple-phase CT scan was performed during the immediate postoperative period to evaluate for early complications, assess the patency of vital structures, and to establish a baseline of the post-ablation bed, as has been previously reported (12, 28, 29). Ablation success was evaluated at 3 months post-IRE treatment via triple-phase CT scan following pancreatic imaging protocol, along with CA19-9, and PET-CT. Ablation success and recurrence have been previously defined (15). Participating institutions standardized utilization of CT scans to avoid the difficulty encountered with cross-comparing CT scans to MRI or CT scan to PET scans in previous studies. Response and progression were evaluated using the international criteria proposed by RECIST 1.1 (21). Serial imaging over at least two months were subsequently used to detect recurrence through study comparison in combination with clinical and serum CA19-9 studies. If equivocal findings were seen on CT then a PET was obtained to either confirm or refute local and/or regional recurrence when required.

Statistical Analyses

OS was defined as the time from the start of treatment to the date of death, due to any reason. PFS was defined as the time from the start of initial IRE treatment to the date of first observed disease progression. The rates of OS and PFS were estimated by Kaplan-Meier method. Multivariable Cox survival regression was performed to determine independent predictors of PFS and OS after backward selection (criterion $p<0.05$) to include all variables of interest. All statistical analyses were performed using SAS version 9.3 (SAS Institute, Cary, NC), and p values less than 0.05 were considered significant.

RESULTS

An intention to treat analysis of 187 patients who met inclusion criteria underwent IRE for stage III LAPC. Baseline demographics of the study cohort are represented in Table 1. Sixty five percent of the cohort was male with a median age of 62 years. The majority of the population were of White (55%) or Asian (39%) ethnicity. Preoperative tumor characteristics and chemotherapy and radiation interventions are represented in Table 2. Thirty eight percent of tumors were located in the head of the pancreas with 53% of patients had tumors > 3 cm in greatest diameter. Preoperative radiation therapy was administered to 28% of the cohort. All patients in this study received induction preoperative chemotherapy. Forty-two patients (22%) within the cohort received FOLFIRINOX alone, 62 (33%) had FOLFIRINOX + gemcitabine and abraxane, and 19 (10%) were administered gemcitabine alone, respectively. A majority of patients (90%) had abnormal CA19- levels at the time of diagnosis once their bilirubin’s were normalized.

Table 3 outlines the operative characteristics, adjunctive procedures, and outcome measures for the cohort. The median time from diagnosis to IRE treatment was 4 months, with 56% of

| Characteristic                      | Study cohort (n=187) |
|-------------------------------------|---------------------|
| Age (years), median, (IQR)          | 62 (21 - 91)        |
| Male gender, n (%)                  | 121 (65)            |
| BMI, median (IQR)                   | 25.7 (14 - 41)      |
| Ethnicity, n (%)                    |                     |
| Asian                               | 73 (39)             |
| Black/African American               | 8 (4)               |
| Native Hawaiian or other Pacific Islander | 1 (0.5)         |
| White                               | 102 (55)            |
| Unknown/not reported                | 2 (1)               |
| Other                               | 1 (0.5)             |
| Past medical history, n (%)         |                     |
| Cardiac                             | 16 (9)              |
| Diabetes                            | 25 (13)             |
| Hypertension                        | 32 (17)             |
| Liver dysfunction                   | 2 (1)               |
| Pancreatitis                        | 12 (6)              |
| Pulmonary                           | 7 (4)               |
| Vascular                            | 4 (2)               |
| Tobacco History                     | 23 (12)             |
| Alcohol Abuse                       | 8 (4)               |
| Past surgical history, n (%)        |                     |
| Appendectomy                        | 13 (7)              |
| Bile Stents                         | 25 (13)             |
| Cholecystectomy                     | 30 (16)             |
| Colon                               | 4 (2)               |
| Distal Pancreatectomy               | 4 (2)               |
| Gastric Bypass                      | 1 (0.5)             |
| Orthopedic                          | 13 (7)              |
| TAH                                 | 10 (5)              |
| Whipple                             | 4 (2)               |
| Karnofsky Performance Score, n (%)  |                     |
| 100%                                | 77 (41)             |
| 90%                                 | 58 (10)             |
| 80%                                 | 47 (4)              |
| 70%                                 | 3 (2)               |
| 0%                                  | 2 (1)               |

IQR, interquartile range; BMI, body mass index; TAH, total abdominal hysterectomy.
the cohort receiving additional adjunctive procedures at the time of IRE. Cholecystectomy (30%) and jejunostomy tube placement (45%) were the most common adjunctive procedures. Thirty-two patients (17%) had local recurrences and 49 (26%) experienced distant recurrence. Mean time to local recurrence was 16.4 months and 15.9 months to distant recurrence from IRE. The liver represented the most common location of distant progression (26%, 13/49). Adverse events following IRE liver represented the most common location of distant progression (26%, 13/49). The patients (17%) had local recurrences and 49 (26%) experienced distant recurrence from IRE. The patients requiring pullback, n (%) 166 (89). Patients with adjunctive procedures 105 (56).

### TABLE 3 | Operative characteristics, adjunctive procedures, and outcome measures.

| Characteristic                                   | Study cohort (n= 187) |
|--------------------------------------------------|-----------------------|
| Time from diagnosis to IRE treatment (mo.), median, (IQR) | 4 (12 – 69) |
| Total IRE delivery time (min.), median, (IQR)     | 49 (2 - 307) |
| Total probe placement time (min.), median (IQR)   | 15 (2 - 21) |
| Total procedure time (min.) median, (IQR)         | 164 (50 - 540) |
| Time from IRE to discharge (days), median, (IQR)  | 7 (5 - 27) |
| Patients requiring pullback, n (%)                | 166 (89) |
| Patients with adjunctive procedures               | 105 (56) |

### Adjunctive Procedure, n (%)

- Cholecystectomy 57 (30)
- Distal pancreatectomy 3 (2)
- Gastrojejunostomy 39 (21)
- Hepaticejunostomy 27 (14)
- Jejunalostomy-tube 85 (45)
- Portal vein or SMV resection 12 (6)
- Subtotal pancreatectomy with celiac resection 5 (3)
- Whipple 17 (9)
- Other (hemia repair, gastro-jejunostomy) 30 (16)

Patients receiving adjuvant therapy during follow-up, n (%) 74 (40). Time to local recurrence from diagnosis, (mo), mean, (IQR) 22.3 (0.1 - 77.1). Time to local recurrence from IRE, (mo), mean, (IQR) 16.4 (0 - 52.5). Time to distant recurrence from diagnosis, (mo), mean, (IQR) 21.9 (0.1 - 90.8). Time to distant recurrence from IRE, (mo), mean, (IQR) 15.9 (0 - 52.5). PFS from diagnosis, (mo), median, (IQR) 21.7 (0.1 - 77.1). PFS from IRE, (mo), median, (IQR) 16.1 (0 - 52.9). OS from diagnosis, (mo), median, (IQR) 25.3 (0.1 - 90.8). OS from IRE, (mo), median, (IQR) 22.4 (0 - 52.5). Recurrence type, n (%)

#### Local

- Bone 1 (0.5)
- Liver 25 (13)
- Lung 13 (7)
- Ascites 2 (1)
- Left suprarenal space 1 (0.5)
- Omentum 1 (0.5)
- Peritoneum 4 (2)
- RUQ small intestine 1 (0.5)
- Regional lymph node disease 1 (0.5)
- Retroperitoneal lymph node 1 (0.5)
- Abdominal wall 1 (0.5)
- Around mesenteric artery 1 (0.5)

#### Distant

- Bone 49 (26)

#### Location of distant progression, n (%)

- Bone 1 (0.5)
- Liver 25 (13)
- Lung 13 (7)
- Ascites 2 (1)
- Left suprarenal space 1 (0.5)
- Omentum 1 (0.5)
- Peritoneum 4 (2)
- RUQ small intestine 1 (0.5)
- Regional lymph node disease 1 (0.5)
- Retroperitoneal lymph node 1 (0.5)
- Abdominal wall 1 (0.5)
- Around mesenteric artery 1 (0.5)

### Supplementary Table 1

#### Operative characteristics, adjunctive procedures, and outcome measures.

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- Total IRE delivery time (min.), median, (IQR) 49 (2 - 307)
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- Patients requiring pullback, n (%) 166 (89)
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#### Adjunctive Procedure, n (%)

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- Whipple 17 (9)
- Other (hemia repair, gastro-jejunostomy) 30 (16)

Patients receiving adjuvant therapy during follow-up, n (%) 74 (40).

#### Local Recurrence type, n (%)

- Bone 1 (0.5)
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- Lung 13 (7)
- Ascites 2 (1)
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- RUQ small intestine 1 (0.5)
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- Retroperitoneal lymph node 1 (0.5)
- Abdominal wall 1 (0.5)
- Around mesenteric artery 1 (0.5)

#### Distant

- Bone 49 (26)

#### Location of distant progression, n (%)

- Bone 1 (0.5)
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- RUQ small intestine 1 (0.5)
- Regional lymph node disease 1 (0.5)
- Retroperitoneal lymph node 1 (0.5)
- Abdominal wall 1 (0.5)
- Around mesenteric artery 1 (0.5)
### TABLE 4 | Risk factors for overall and progression free survival in stage III locally advanced pancreatic cancer patients.

| Characteristic                        | KM Median (95% CI) | P value |
|---------------------------------------|--------------------|---------|
| **OS from diagnosis**                 |                    |         |
| CA19-9 change from diagnosis to IRE   |                    |         |
| Abnormal to normal                    | 30.2 (22.5-43.5)   | 0.03    |
| Abnormal to abnormal                  | 18.8 (15.4-22.7)   |         |
| Vascular involvement                  |                    |         |
| ≤ 180°                                | 52.8 (17.7-90.8)   | 0.006   |
| > 180°                                | 22.7 (18.4-24.7)   |         |
| Prior chemotherapy duration           |                    |         |
| > 5 months                            | 23.1 (17.4-34.7)   | 0.04    |
| ≤ 5 months                            | 21.7 (15.7-24.4)   |         |
| **OS from IRE**                       |                    |         |
| Age                                   |                    |         |
| ≤ 61                                 | 23.9 (10.4-47.7)   | 0.02    |
| > 61                                 | 18.2 (13.3-23.1)   |         |
| CA19-9 change from diagnosis to IRE   |                    |         |
| Normal at IRE (≤ 37 U/mL)             | 17.5 (12.4-22.8)   | 0.02    |
| Abnormal at IRE (> 37 U/mL)           | 9.3 (5.9-13.3)     |         |
| Diabetes                              |                    |         |
| Yes                                  | 13.3 (8.4-14.8)    | 0.03    |
| No                                   | 23.1 (22.0-31.2)   |         |
| Vascular involvement                  |                    |         |
| ≤ 180°                               | 52.5 (4.8-62.5)    | 0.01    |
| > 180°                               | 12.4 (6.6-16.4)    |         |
| Number of comorbidities               |                    |         |
| ≤ 2                                  | 21.6 (10.3-23.9)   | 0.04    |
| > 2                                  | 12.4 (6.6-16.4)    |         |
| Prior chemotherapy                    |                    |         |
| FOLFIRINOX + gemcitabine/abraxane     | 23.2 (19.4-35.9)   | 0.01    |
| FOLFIRINOX                            | 13.3 (9.4-18.2)    |         |
| Gemcitabine/abraxane                  | 12.4 (4.6-26.5)    |         |
| Prior radiation                       |                    |         |
| Yes                                  | 12.3 (8.4-17.2)    |         |
| No                                   | 26.5 (22.7-63.6)   |         |
| Tumor size                            |                    |         |
| ≤ 3.6                                | 26.5 (22.5-2.5)    | 0.0003  |
| > 3.6                                | 18.2 (15.8-22.8)   |         |
| **PFS from diagnosis**                |                    |         |
| Vascular involvement                  |                    |         |
| ≤ 180°                               | 52.8 (17.5-77.1)   | 0.001   |
| > 180°                               | 16.8 (15.7-18.4)   |         |
| Prior chemotherapy                    |                    |         |
| FOLFIRINOX + Gemcitabine/abraxane     | 20.8 (17.5-24.7)   | 0.04    |
| FOLFIRINOX                            | 18.7 (16.3-20.7)   |         |
| Gemcitabine/abraxane                  | 15.5 (10.6-25.4)   |         |
| **PFS from IRE**                      |                    |         |
| Age                                   |                    |         |
| ≤ 61                                 | 22.0 (18.8-30.4)   | 0.046   |
| > 61                                 | 12.5 (8.9-18.6)    |         |
| CA19-9                                |                    |         |
| Normal at IRE                         | 8.9 (7.0-11.5)     | 0.006   |
| Abnormal at IRE                       | 5.3 (3.7-6.6)      |         |
| Diabetes                              |                    |         |
| Yes                                  | 7.2 (4.3-13.8)     | 0.0003  |
| No                                   | 20.6 (14.6-22.8)   |         |
| Vascular involvement                  |                    |         |
| ≤ 180°                               | 24.2 (4.5-52.5)    | 0.005   |
| > 180°                               | 7.3 (5.8-8.8)      |         |
| Prior chemotherapy                    |                    |         |
| FOLFIRINOX + gemcitabine/abraxane     | 19.4 (14.5-23.2)   | <0.0001 |
| FOLFIRINOX                            | 8.7 (5.8-11.5)     |         |
| Gemcitabine/abraxane                  | 8.8 (2.9-10.9)     |         |

(Continued)

### TABLE 4 | Continued

| Characteristic                        | KM Median (95% CI) | P value |
|---------------------------------------|--------------------|---------|
| **Preoperative radiation**            |                    | <0.0001 |
| Yes                                  | 6.7 (3.3-8.9)      |         |
| No                                   | 22.3 (18.9-29.3)   |         |
| **Tumor size (cm)**                   |                    | <0.0001 |
| ≤ 3.6                                | 22.8 (19.4-35.9)   |         |
| > 3.6                                | 10.4 (8.2-16.1)    |         |

OS, overall survival; KM, Kaplan Meier; CA19-9, cancer antigen 19-9; IRE, irreversible electroporation; PFS, progression free survival; FOLFIRINOX, folinic acid, 5-fluorouracil, infuscin, and oxaplatin.

Bold values represent statistical significance with p values less than 0.05.

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Monitoring of response to neoadjuvant treatment by preoperative CA 19-9 levels was a significant factor in predicting survival of LAPC by IRE. OS from diagnosis and from IRE was significantly increased when patients with abnormally elevated CA19-9 levels reached normative values (<37U/mL) at the time of operation (30.2 mo. vs. 18.8 mo., 95% CI, 22.5 – 43.5, p=0.03), (17.5 mo. vs. 9.3 mo., 95% CI, 12.4 – 22.8, p =0.02). This finding was replicated on PFS measured from the time of surgery when CA19-9 reached normal levels at the time of IRE (8.9 mo. vs 5.3 mo., p=0.006) (Table 4).

On univariate Kaplan Meier (KM) analyses when the 25 patients who underwent pancreatectomy with IRE were compared two patients who underwent IRE alone there was a similar median overall progression free survival from diagnosis with margin accentuation 21.4 (11.1 two 24.3 parentheses versus insight 2 22.6 (19.9 two 25.4) months, p=0.0690, and a statistically significant improvement and overall progression free survival from IRE treatment pancreatectomy with IRE 10.2 (3.4 to 31.5) versus in-situ 21.9 (16.1 to 23.3) months, p=0.0452 (Supplemental Figure 1). Overall survival from diagnosis for pancreatectomy with IRE was 24.9 (12.6 to 39.1) versus 29.4 (23.1 to 36.2) months, p=0.23 and from IRE treatment was 15.6 mon (7.6 to 33.6) vs IRE treatment was 15.6 mon (7.6 to 33.6) months, p=0.076.

On multivariable Cox regression analysis for independent predictive factors for survival, (Table 5) abnormal CA19-9 values at IRE (HR 2.16, 95% CI 1.1-4.2, p=0.02), and chemotherapy duration ≤ 5 months (HR 1.98, 95% CI 1.02-3.86, p=0.04) independently predicted a worse OS from diagnosis (Figures 1 A, B). However, age ≤ 61 years (HR 0.4, 95% CI, 0.21-0.78, p=0.008) and those without prior radiation history (HR 0.49, 95% CI 0.26-0.93, p=0.03) were independent predictors of improved in OS form IRE. Age ≤ 61 (HR 0.53, 95% CI 0.28-0.99, p=0.046) and FOLFIRINOX plus gemcitabine/abraxane induction...
The most important finding from this study is the identification of clinicopathologic characteristics that predict survival following open in-situ IRE for LAPC, which has not been previously established. Significant progress in oncologic management of stage III LAPC has occurred in the past decade (30). Total neoadjuvant chemotherapy in the setting of metastatic and locoregional pancreatic adenocarcinoma has improved survival and allowed for more aggressive and consolidative operative interventions (5, 31–33). IRE in the setting of LAPC is one example and was first described in 2012, and has been proven safe near vital vessels and ductal structures due to its non-thermal mechanism of action (7). IRE has further proven to be an effective palliative surgical intervention with remarkable improvements in OS and PFS for those diagnosed with LAPC (8, 12, 34). Therefore, a better understanding of preoperative factors to assist in selecting patients to undergo IRE is now critical with the establishment of these key outcome measures.

The median PFS of 21.7 months and OS of 25.5 months in these 187 LAPC patients treated with open in-situ IRE, confirms that of previous reports and warranted this investigation into the selection process within this registry (8, 11, 12, 34–36). Earlier publications demonstrated variation in OS when utilizing IRE for LAPC (37–39). However, many factors may explain this underlying discrepancy. IRE has a demonstrable learning curve and over time with performance of more cases, allows for completion of complex ablations involving larger tumors and those with a high degree of vascular involvement (40). Differences in tumor biology, heterogeneity of NAC regimens, patient selection, and approach or technique (open vs laparoscopic vs percutaneous) technique may also attribute. Lack of energy delivery standardization also greatly limits the reproducibility of results. Inadequate energy delivery to the tumor leading to incomplete ablations or reversible electroperoration can actually increase tumor growth (26, 41).

Differences in tumor biology, heterogeneity of NAC regimens, patient frailty scores, which is a known independent predictor of mortality over time. Elderly patients are also more likely to have increased frailty scores, which is a known independent predictor of mortality following pancreaticoduodenectomy (48–50). It should be noted that of previous reports and warranted this investigation into the selection process within this registry (8, 11, 12, 34). However, many factors may explain this underlying discrepancy. IRE has a demonstrable learning curve and over time with performance of more cases, allows for completion of complex ablations involving larger tumors and those with a high degree of vascular involvement (40). Differences in tumor biology, heterogeneity of NAC regimens, patient selection, and approach or technique (open vs laparoscopic vs percutaneous) technique may also attribute. Lack of energy delivery standardization also greatly limits the reproducibility of results. Inadequate energy delivery to the tumor leading to incomplete ablations or reversible electroperoration can actually increase tumor growth (26, 41).

Here we observed age > 61, > 2 comorbidities, and those with diabetes to negatively impact OS and PFS following IRE. These findings are consistent with reports seen in other pancreatic adenocarcinoma patient populations (4, 42–44). Age less than 61 at electroporation also independently predicted prolonged PFS and OS following IRE in this cohort. It is well known that the incidence of PDAC positively correlates with age (45). Several population-based studies have reported on poorer prognoses in older PDAC patients (46, 47). Wang et al. recently reported on 126,066 patients from the National Cancer Institute’s Surveillance, Epidemiology, and the End Results data base. Risk for mortality was double for those aged 40–80 years compared to PDAC patients less than 40 years (42). These findings are expected as PDAC is primarily a cancer of older age and physiologic reserve decreases over time. Elderly patients are also more likely to have increased frailty scores, which is a known independent predictor of mortality following pancreaticoduodenectomy (48–50). It should be noted that 41% of this cohort had performance statuses of 100%, which demonstrates our selection bias toward optimal function prior to operative intervention. The importance of performance status (PS) as a prognosticator in all stages of PDAC cannot be overstated (51). Every effort should be taken to optimize PS for LAPC patients presenting for IRE as nearly every patient will have undergone extensive induction chemotherapy leaving them

TABLE 5 | Independent risk factors for survival in locally advanced pancreatic cancer patients.

| OS from diagnosis | Adjusted HR (95% CI) | P value |
|-------------------|---------------------|---------|
| CA19-9 change from diagnosis to IRE | Abnormal to abnormal | 2.159 (1.1-4.2) | 0.02 |
| Age | ≤ 61 | 0.4 (0.21-0.78) | 0.008 |
| Prior radiation | No | 0.49 (0.26-0.93) | 0.03 |
| PFS from IRE | Abnormal to normal | 2.19 (1.14-4.201) | 0.018 |
| Induction chemotherapy | FOLFIRINOX + gemcitabine/abraxane | 0.37 (0.15-0.89) | 0.027 |
| | Gemcitabine/abraxane | 0.48 (0.28-0.82) | 0.03 |

OS, overall survival; HR, hazard ratio; CI, confidence interval; CA19-9, cancer antigen 19-9; IRE, irreversible electroporation; FOLFIRINOX, folinic acid, 5-fluorouracil, irinotecan, and oxaliplatin.

**DISCUSSION**

The most important finding from this study is the identification of clinicopathologic characteristics that predict survival following open in-situ IRE for LAPC, which has not been previously established. Significant progress in oncologic management of stage III LAPC has occurred in the past decade (30). Total neoadjuvant chemotherapy in the setting of metastatic and locoregional pancreatic adenocarcinoma has improved survival and allowed for more aggressive and consolidative operative interventions (5, 31–33). IRE in the setting of LAPC is one example and was first described in 2012, and has been proven safe near vital vessels and ductal structures due to its non-thermal mechanism of action (7). IRE has further proven to be an effective palliative surgical intervention with remarkable improvements in OS and PFS for those diagnosed with LAPC (8, 12, 34). Therefore, a better understanding of preoperative factors to assist in selecting patients to undergo IRE is now critical with the establishment of these key outcome measures.

The median PFS of 21.7 months and OS of 25.5 months in these 187 LAPC patients treated with open in-situ IRE, confirms that of previous reports and warranted this investigation into the selection process within this registry (8, 11, 12, 34–36). Earlier publications demonstrated variation in OS when utilizing IRE for LAPC (37–39). However, many factors may explain this underlying discrepancy. IRE has a demonstrable learning curve and over time with performance of more cases, allows for completion of complex ablations involving larger tumors and those with a high degree of vascular involvement (40). Differences in tumor biology, heterogeneity of NAC regimens, patient selection, and approach or technique (open vs laparoscopic vs percutaneous) technique may also attribute. Lack of energy delivery standardization also greatly limits the reproducibility of results. Inadequate energy delivery to the tumor leading to incomplete ablations or reversible electroperoration can actually increase tumor growth (26, 41).

Differences in tumor biology, heterogeneity of NAC regimens, patient frailty scores, which is a known independent predictor of mortality over time. Elderly patients are also more likely to have increased frailty scores, which is a known independent predictor of mortality following pancreaticoduodenectomy (48–50). It should be noted that 41% of this cohort had performance statuses of 100%, which demonstrates our selection bias toward optimal function prior to operative intervention. The importance of performance status (PS) as a prognosticator in all stages of PDAC cannot be overstated (51). Every effort should be taken to optimize PS for LAPC patients presenting for IRE as nearly every patient will have undergone extensive induction chemotherapy leaving them
malnourished and immune compromised. Prescribing preoperative nutritional supplements is one way providers can positively influence post-IRE outcomes (52).

In regard to comorbidities, many studies have established risk for the development of PDAC in the setting of diabetes mellitus (DM) (53). Additionally, our finding of DM negatively impacting OS (13.3 mo. vs. 23.1 mo., p<0.003) and PFS (20.6 mo. vs. 7.2 mo., p<0.0003) following IRE is in agreement with current knowledge (54–57). The effect of DM on survival has been demonstrated in both the short- and long-term settings. In 2015 Yuan et al., demonstrated significantly decreased OS in PDAC patients diagnosed with long term (>4 years) DM (58). Chu et al. also reported recent onset DM as an independent predictor of post resection survival (43). DM has also been significantly associated with increased tumor sizes and increased risk of death following pancreatectomy and adjuvant chemotherapy (59). Collectively, these data suggest at this time patients with DM are poor candidates to receive IRE for LAPC and warrants further investigations into treatment options for such an at risk population.

Anatomic tumor characteristics are also important to consider when evaluating patients for IRE therapy. Survival among LAPC patients following NAC and resection has been integrally tied to tumor size. Gemenetiz et al. found significantly prolonged OS in resected patients with smaller tumor sizes (33). Smaller tumors were also independently predictive of survival in an ancillary study of the LAP 07 trial (60). In agreement with these assessments, we have also found smaller tumor size

![FIGURE 1](https://example.com/figure1.png)

**FIGURE 1** | Independent predictors of overall survival from diagnosis. (A) Overall survival comparison by change in CA19-9 status from diagnosis to IRE. (B) Overall survival comparison by induction chemotherapy duration.
(<3.6cm) to be predictive of PFS and OS following IRE on univariate analyses. In addition, patients with vascular involvement ≤180 degrees of their affected structure are more likely to have significantly longer OS from their initial diagnoses. Thus, it appears IRE may have its greatest impact on tumors of smaller size with less circumferential vessel involvement. Surgeons and interventionalists performing IRE should be cognizant of these tumor qualities and discuss in such cases in a multidisciplinary setting prior to proceeding.

This data again highlights the important prognostic value of CA19-9 in LAPC and adds value to its ability to be used as a treatment biomarker. Serum measurement of CA19-9 as a surrogate for clinical outcomes in pancreatic cancer is well established (61–63). However, only recently have we begun to accept CA19-9 monitoring to guide multimodality therapy. Following NAC, normalization of CA19-9 has been reported to be a strong prognostic marker for long term survival (64–66). Here, patients who achieved normative CA19-9 levels at the time of IRE were able to achieve nearly double the survivals of those with abnormal values. This reiterates that CA19-9 should be used to guide multimodality therapy and suggests those with good response may be better candidates for further consolidative therapy.

The effectiveness of IRE as a consolidative therapy in conjunction with systemic chemotherapy and/or chemoradiation is becoming better understood. However, current NCCN guidelines continue to be heterogeneous in chemotherapeutic recommendations for stage III LAPC (67). This array of treatment options may allow for more tolerable treatment to be prescribed yet with limited efficacy that may confound key outcomes. The success of FOLFIRINOX in treatment of LAPC or borderline resectable disease calls for more standardization of preoperative treatment (5, 68, 69). Our finding that FOLFIRINOX
utility in LAPC is fraught with controversy, yet guidelines recommend its use. Though, data surrounding its use has historically been controversial, however current NCCN guidelines are likely underestimated. Additionally, there was a lack of randomization in this prospective cohort limits ability to determine the degree of recurrence or progressive disease based on current RECIST guidelines are likely underestimated. Furthermore, the delivery of electroporation with these chemotherapeutics (i.e. electrochemotherapy) improves the delivery of such agents to a complex tumor microenvironment (TME) with synergistic anti-tumor activity (70, 71). With respect to radiation, use of radiotherapy in the setting of LAPC has been described (76). However, at this time as an explanation of these findings would be speculative at best. Certainly, more research is needed to investigate the impact radiation has in the setting of IRE and the interplay of their mechanisms of action within the TME.

IRE has been described as a last resort, with some practitioners referring for intervention once patients have become otherwise unresponsive to systemic chemotherapy. For example, Piella et al. reported on 10 patients with unresponsive LAPC who underwent IRE and found median OS of 7.5 months (38). In light of the present findings, we want to strongly encourage the medical oncology community to refer for IRE in patients whose biology of disease and clinical characteristics would positively favor their response to IRE. To that end, we have recommended a specific treatment algorithm that optimizes all three active treatments in the management of LAPC (Figure 3). We believe it is critical to understand that potential substantial benefit can be obtained when all of these favorable prognostic factors are achieved, but we also want to emphasize that improvements in overall outcomes can be achieved even in patients who may not meet all of these prognostic factors and represents a guide for patient selection and for future management and comparisons.

The present study should be interpreted with respect to the following limitations. First, lack of randomization in this prospective cohort limits ability to determine the degree of survival benefit patients received from induction chemotherapy and IRE. Conventional imaging modalities to detect immune relevant responses are lacking and therefore determination of recurrence or progressive disease based on current RECIST guidelines are likely underestimated. Additionally, there was a degree of post IRE imaging variability between participating institutions. As previously reported, this cohort is prone to selection bias. The participating centers have carefully selected patients who do not progress on systemic chemotherapy, with enhanced performance statuses, and limited co-morbidities to receive IRE. These limitations notwithstanding, this study is the most comprehensive and only prospective multi-institution evaluation for prognosticators of survival in the setting of LAPC treated with open technique IRE to date. Until now, the optimal patient characteristics highlighting improved OS and PFS after IRE for LAPC were not elucidated.

**CONCLUSIONS**

This prospective cohort evaluation of stage III LAPC patients treated with open IRE demonstrates prominent factors predictive of PFS and OS that should be used to aide in selection or referral for patients to receive open technique IRE. These results demonstrate that prolonged survival beyond historical controls can be achieved by IRE of LAPC in appropriately selected patients. This study further supports the design of randomized multi-center trials investigating the efficacy of IRE, which are now actively recruiting participants (NCT03899636, NCT03899649).

**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.
ETHICS STATEMENT

The studies involving human participants were reviewed and approved by U of L IRB. The patients/participants provided their written informed consent to participate in this study.

AUTHOR CONTRIBUTIONS

MRW, KW, EK, MJW, JC, RW, and RMII contributed to study design and manuscript revisions. MRW, KW, and RMII contributed to study design, drafting of the manuscript, and manuscript revisions. EK, MJW, JC, RW, and RMII contributed to study design, data acquisition, and manuscript revisions. RM contributed to study design, data acquisition, analysis, drafting of the manuscript, and manuscript revisions. All authors contributed to the article and approved the submitted version.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fonc.2021.817220/supplementary-material

Supplementary Figure 1 | Progression free survival of patients who underwent Pancreatectomy with IRE for margin accentuation versus patients who underwent IRE alone (Ir-Stu)

Supplementary Table 1 | Observed adverse events.
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Conflict of Interest: RM II, MD, PhD, FACS is an educational consultant for AngioDynamics, Inc.

The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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