## UNIVERSITYOF BIRMINGHAM

# University of Birmingham Research at Birmingham

## **Adjuvant Chemotherapy Associated with Survival** Benefit Following Neoadjuvant Chemotherapy and **Pancreatectomy for Pancreatic Ductal** Adenocarcinoma

Kamarajah, Sivesh K; White, Steven A; Naffouje, Samer A; Salti, George I; Dahdaleh, Fadi

DOI:

10.1245/s10434-021-09823-0

License:

Creative Commons: Attribution (CC BY)

Document Version

Publisher's PDF, also known as Version of record

Citation for published version (Harvard):

Kamarajah, SK, White, SA, Naffouje, SA, Salti, GI & Dahdaleh, F 2021, 'Adjuvant Chemotherapy Associated with Survival Benefit Following Neoadjuvant Chemotherapy and Pancreatic Ductal Adenocarcinoma: A Population-Based Cohort Study', Annals of Surgical Oncology, vol. 28, no. 11, pp. 6790-6802. https://doi.org/10.1245/s10434-021-09823-0

Link to publication on Research at Birmingham portal

**General rights** 

Unless a licence is specified above, all rights (including copyright and moral rights) in this document are retained by the authors and/or the copyright holders. The express permission of the copyright holder must be obtained for any use of this material other than for purposes

- •Users may freely distribute the URL that is used to identify this publication.
- •Users may download and/or print one copy of the publication from the University of Birmingham research portal for the purpose of private study or non-commercial research.
- •User may use extracts from the document in line with the concept of 'fair dealing' under the Copyright, Designs and Patents Act 1988 (?)
  •Users may not further distribute the material nor use it for the purposes of commercial gain.

Where a licence is displayed above, please note the terms and conditions of the licence govern your use of this document.

When citing, please reference the published version.

While the University of Birmingham exercises care and attention in making items available there are rare occasions when an item has been uploaded in error or has been deemed to be commercially or otherwise sensitive.

If you believe that this is the case for this document, please contact UBIRA@lists.bham.ac.uk providing details and we will remove access to the work immediately and investigate.

Download date: 16. Sep. 2023

#### ORIGINAL ARTICLE - HEPATOBILIARY TUMORS

### Adjuvant Chemotherapy Associated with Survival Benefit Following Neoadjuvant Chemotherapy and Pancreatectomy for Pancreatic Ductal Adenocarcinoma: A Population-Based **Cohort Study**

Sivesh K. Kamarajah, BMedSci, MBChB<sup>1,2,3</sup>, Steven A. White, MD, FRCS<sup>1,2</sup>, Samer A. Naffouje, MD<sup>4</sup>, George I Salti, MD<sup>5,6</sup>, and Fadi Dahdaleh, MD<sup>6</sup>

<sup>1</sup>Department of HPB and Transplant Surgery, The Freeman Hospital, Newcastle upon Tyne, Tyne and Wear, UK; <sup>2</sup>Institute of Cellular Medicine, Newcastle University, Newcastle Upon Tyne, Newcastle, UK: <sup>3</sup>Department of Surgery, Oueen Elizabeth Hospital Birmingham, University Hospital Birmingham NHS Trust, Birmingham, UK; <sup>4</sup>Department of Surgical Oncology, H. Lee Moffitt Cancer and Research Institute, Tampa, FL; <sup>5</sup>Department of General Surgery, University of Illinois Hospital and Health Sciences System, Chicago, IL; <sup>6</sup>Department of Surgical Oncology, Edward-Elmhurst Health, Naperville, IL

#### ABSTRACT

**Background.** Data supporting the routine use of adjuvant chemotherapy (AC) compared with no AC (noAC) following neoadjuvant chemotherapy (NAC) and resection for pancreatic ductal adenocarcinoma (PDAC) are lacking. This study aimed to determine whether AC improves longterm survival in patients receiving NAC and resection.

Methods. Patients receiving resection for PDAC following NAC from 2004 to 2016 were identified from the National Cancer Data Base (NCDB). Patients with a survival rate of < 6 months were excluded to account for immortal time bias. Propensity score matching (PSM) and Cox regression analysis were performed to account for selection bias and analyze the impact of AC on overall survival.

**Results.** Of 4449 (68%) noAC patients and 2111 (32%) AC patients, 2016 noAC patients and 2016 AC patients

**Supplementary Information** The online version contains supplementary material available at https://doi.org/10.1245/s10434-

© The Author(s) 2021

First Received: 6 October 2020 Accepted: 26 February 2021

F. Dahdaleh, MD

e-mail: Fadi.Dahdaleh@EEHealth.org

021-09823-0

chemotherapy (AC) or chemoradiotherapy 1-3 remains the standard treatment for localized pancreatic ductal adenocarcinoma (PDAC). Recently, neoadjuvant therapy (NAT) for PDAC is recognized as an acceptable treatment strategy for patients with resectable, borderline resectable, and locally advanced disease. 4-7 Neoadjuvant chemotherapy

(NAC) and/or neoadjuvant radiotherapy (NART) help to

Multimodal therapy with pancreatectomy and adjuvant

remained after PSM. After matching, AC was associated

with improved survival (median 29.4 vs. 24.9 months;

p < 0.001), which remained after multivariable adjustment (HR 0.81, 95% confidence interval [CI] 0.75-0.88;

p < 0.001). On multivariable interaction analyses, this

benefit persisted irrespective of nodal status: N0 (hazard ratio [HR] 0.80, 95% CI 0.72–0.90; p < 0.001), N1 (HR

0.76, 95% CI 0.67-0.86; p < 0.001), R0 margin status (HR

0.82, 95% CI 0.75–0.89; p < 0.001), R1 margin status (HR 0.77, 95% CI 0.64–0.93; p = 0.007), no neoadjuvant

radiotherapy (NART; HR 0.84, 95% CI 0.74-0.96;

p = 0.009), and use of NART (HR 0.80, 95% CI 0.73–0.88;

p < 0.001). Stratified analysis by nodal, margin, and

Conclusion. AC following NAC and resection is associ-

ated with improved survival, even in margin-negative and

node-negative disease. These findings suggest completing planned systemic treatment should be considered in all

NART status demonstrated consistent results.

resected PDACs previously treated with NAC.

Published online: 30 March 2021

downstage the size and anatomic extent of the primary tumor, to improve the likelihood of a margin-negative (R0) resection, and improve the selection of patients with biologically aggressive cancers who are less likely to benefit from surgery. <sup>8,9</sup> Furthermore, NAT may also maximize the number of patients receiving multimodal therapy, as up to 40% of patients do not receive AC after pancreatectomy. <sup>8,9</sup> Indeed, administering all planned systemic therapy preoperatively may obviate the need for additional AC following resection, but accumulated toxicity from NAT may theoretically prohibit curative-intent surgery. Therefore, most NAT protocols include only a portion of total planned cycles to be administered upfront. <sup>10–12</sup>

Although the benefit of AC to patients who have undergone resection first is clear, the benefit of AC after NAC and resection is less well-established. First, previous retrospective analyses investigating the added benefit of AC after NAT and resection have yielded conflicting results, as some suggested benefit in certain subgroups only. Second, while guidelines from the American Society of Clinical Oncology (ASCO) and National Comprehensive Cancer Network (NCCN) recommend considering additional AC in this setting, those recommendations likely represent an extrapolation from data of adjuvant trials. Therefore, more high-quality data are needed to truly understand the impact of AC following NAT in patients undergoing resection for PDAC.

We sought to add evidence to this debate by performing a large, nationwide retrospective study to assess the potential benefit of AC after NAC and resection of PDAC. With contemporary data from the National Cancer Data Base (NCDB), we analyzed the association of AC with survival after resection of PDAC in patients surviving > 6 months to account for immortal time bias. We also used propensity score matching (PSM) to address treatment selection bias and assessed survival in clinically relevant subgroups of patients based on nodal and margin status.

#### **METHODS**

Data Source

The NCDB is a joint project of the Commission on Cancer (CoC) of the American College of Surgeons and the American Cancer Society. <sup>20,21</sup> Data from over 1500 CoCaccredited hospitals are gathered to include > 70% of all newly diagnosed cancers in the US. Details on demographics, facility type and location, clinicopathologic characteristics, treatment, and outcomes are available.

Study Population

NCDB was used to identify all patients > 18 years of age diagnosed with non-metastatic PDAC undergoing resection [i.e. pancreaticoduodenectomy (PD) or distal pancreatectomy (DP)] with curative intent between 2004 and 2016. The International Classification of Diseases for Oncology, Third Edition (ICD-O-3) was used to select adenocarcinoma and to exclude other histologies (ICD-O-3 morphology codes 8240–8248). Patients with concomitant cancer diagnoses and those with missing data on receipt of perioperative chemotherapy were excluded. Patients with survival > 6 months were only included to account for immortal time bias in patients who were not able to complete AC.<sup>22</sup>

Center volume is defined as the annual resection volume and was divided into five quintiles (i.e. quintiles 1–5). The following patient-level characteristics were analyzed: age (36–50, 51–65, 66–80, > 80 years), race (White, Black, other), Charlson–Deyo comorbidity score (CDCC), year of diagnosis, insurance status (Medicaid/Medicare, private insurance, uninsured), zip code-level education status (i.e. < 7.0%, 7.0–12.9%, 13.0–20.9%, > 21.0%), nodal status (N0, N1, N2), tumor grade/differentiation (well, moderate, poor, anaplastic), and lymphovascular invasion (absent, present). The 8th edition of the American Joint Committee on Cancer (AJCC) staging system was used both T and N classifications. Finally, we analyzed the rates of receipt of AC as the primary exposure variable.

Statistical Analysis

Categorical variables were compared using the Chisquare test, and non-normally distributed data were analyzed using the Mann-Whitney U test. Survival was estimated using Kaplan-Meier survival curves and compared using the log-rank test. Multivariable analyses used Cox proportional hazards models. The conditional probability of receiving AC, i.e. the propensity score, was estimated using a multivariable logistic regression model including all variables listed above. We then created balanced cohorts using 1-to-1 nearest-neighbor PSM without replacement (caliper width 0.1 standard deviations).<sup>23</sup> Balance diagnostics were conducted using standardized mean differences, with a value < 0.1 indicating good balance.<sup>23</sup> Sensitivity and interaction analyses were performed by nodal status (i.e. N0, N1, N2/3), margin status (i.e. R0, R1), and receipt of NART on long-term survival. A p-value of < 0.05 was considered statistically significant. Data analysis was performed using R Foundation statistical software (R 3.2.2) with TableOne, ggplot2, Hmisc, Matchit, and survival packages (The R Foundation for Statistical Computing, Vienna, Austria) as previously

described.<sup>24,25</sup> This study was exempt from Institutional Review Board approval.

#### **RESULTS**

Clinicopathologic Characteristics and Propensity Score Matching

This study included 6560 patients undergoing surgical resection following NAC for PDAC, of whom 2111 (32%) received AC and 4449 (68%) did not. Baseline characteristics of the entire unmatched cohort are presented in Table 1. Baseline demographics of the unmatched cohort revealed that patients receiving AC were younger and had lower CDCC scores (Table 1). Moreover, AC patients more often had advanced tumor stages and positive lymph nodes. Of patients receiving AC, 60% (1266/2111) had also received NART, compared with 65% in no AC (noAC) patients (p < 0.001). To account for potential treatment selection bias, PSM was performed as described above, which resulted in well-balanced cohorts (Table 1). Standardized mean differences were calculated for each variable and ranged between 0.01 and 0.05, indicating good balance.

Association of Adjuvant Chemotherapy with Survival

For the overall cohort, median survival was 27.1 months and 5-year survival was 23%. In the unmatched cohort, survival of patients receiving AC was significantly longer than those who did not receive AC (median 29.5 vs. 25.9 months; 5-year: 24% vs. 23%; p < 0.013) (Fig. 1, Table 2). After matching, this survival advantage persisted (median 29.4 vs. 24.9 months; 5-year: 24% vs. 20%; p < 0.001) (Fig. 1b, Table 2) and remained after multivariable adjustment [hazard ratio (HR) 0.81, 95% CI 0.75–0.88; p < 0.001] (Tables 2 and 3). On multivariable analysis and after PSM, factors associated with adverse survival included poor education level and median income, single-agent NAC, poor tumor grade, node-positive disease, margin-positive resection, and the presence of lymphovascular invasion (Table 3).

Interaction Between Adjuvant Chemotherapy and Nodal Status

Interaction analyses were performed to further understand the impact of AC by nodal status. In unadjusted analysis, there were significant differences in survival between AC and noAC patients in patients with N1 disease (median 39.1 vs. 34.4 months; p = 0.014) (Fig. 2b), but not N2/N3 disease (median 90.0 vs. 86.1 months; p = 0.1)

(Fig. 2c). On multivariable analyses modeling the interaction between receipt of AC and nodal status, a survival benefit was again seen for patients with N0 and N1 disease but not N2/N3 disease (Table 4, electronic supplementary Table 2). As a sensitivity analysis, three separate multivariable analyses in cohorts including only those with N0, N1, and N2/N3 disease were conducted, respectively. These analyses confirmed the same findings (Table 3).

Interaction Between Adjuvant Chemotherapy and Margin Status

Interaction analyses were performed to further understand the impact of AC by margin status. In unadjusted analysis, there were significant differences in survival between AC and noAC patients in those with R0 disease (median 31.2 vs. 26.7 months; p < 0.001) (Fig. 3a) and in patients with R1 disease (median 22.1 vs. 18.1 months; p = 0.007) (Fig. 3b). On multivariable analyses modeling the interaction between receipt of AC and margin status, a survival benefit was again seen for patients with R0 margin status (HR 0.83, 95% CI 0.76–0.90; p < 0.001) and R1 margin status (HR 0.73, 95% CI 0.57–0.93; p < 0.001) (Table 4, electronic supplementary Table 3). As a sensitivity analysis, we performed two separate multivariable analyses in cohorts including only those with R0 or R1 margin, respectively. These analyses confirmed the same findings (Table 2).

Association of Adjuvant Chemotherapy and Neoadjuvant Radiotherapy with Survival

Additional analyses were performed to further understand the impact of AC in patients who received NART. In unadjusted analysis, there were significant differences in survival between AC and noAC patients in those who did not receive NART (median 31.4 vs. 24.7 months; p = 0.009) (electronic supplementary Fig. 1a) and those who did (median 28.9 vs. 24.9 months; p < 0.001) (electronic supplementary Fig. 1b). On multivariable analyses modeling the interaction between receipt of AC and radiotherapy, a survival benefit was again seen for patients without NART (HR 0.84, 95% CI 0.74–0.96; p < 0.001) and those with NART (HR 0.80, 95% CI 0.73-0.88; p < 0.001) (electronic supplementary Table 4). As a sensitivity analysis, we performed two separate multivariable analyses in cohorts including only those without and with adjuvant radiotherapy, respectively. These analyses confirmed the same findings (Table 2).

TABLE 1 Clinicopathologic characteristics by receipt of adjuvant chemotherapy following neoadjuvant chemotherapy and resection of pancreatic ductal adenocarcinoma in unmatched and matched cohorts

	Unmatched cohort	Unmatched cohort			Matched cohort		
	noAC $[n = 4449]$	AC $[n = 2111]$	p value	noAC $[n = 2061]$	AC $[n = 2061]$	p value	
Center volume							
1 (lowest)	431 (9.7)	290 (13.7)	< 0.001	277 (13.4)	277 (13.4)	1.0	
2	527 (11.8)	278 (13.2)		262 (12.7)	271 (13.1)		
3	704 (15.8)	268 (12.7)		259 (12.6)	264 (12.8)		
4	1257 (28.3)	580 (27.5)		565 (27.4)	571 (27.7)		
5 (highest)	1530 (34.4)	695 (32.9)		698 (33.9)	678 (32.9)		
Facility type							
Community	906 (20.4)	559 (26.5)	< 0.001	544 (26.4)	539 (26.2)	0.8	
Academic	2941 (66.1)	1354 (64.1)		1336 (64.8)	1328 (64.4)		
Others	602 (13.5)	198 (9.4)		181 (8.8)	194 (9.4)		
Facility location							
Northeast	1032 (23.2)	592 (28.0)	< 0.001	560 (27.2)	574 (27.9)	0.9	
South	1598 (35.9)	628 (29.7)		639 (31.0)	620 (30.1)		
Midwest	1240 (27.9)	582 (27.6)		552 (26.8)	571 (27.7)		
West	523 (11.8)	291 (13.8)		295 (14.3)	280 (13.6)		
Unknown	56 (1.3)	18 (0.9)		15 (0.7)	16 (0.8)		
Hospital distance, miles	3						
< 12.5	1536 (34.5)	772 (36.6)	< 0.001	749 (36.3)	754 (36.6)	1.0	
12.5-49.9	1517 (34.1)	788 (37.3)		764 (37.1)	767 (37.2)		
≥ 50 miles	1396 (31.4)	551 (26.1)		548 (26.6)	540 (26.2)		
Year of diagnosis							
2004-2005	214 (4.8)	27 (1.3)	< 0.001	29 (1.4)	27 (1.3)	1.0	
2006-2007	240 (5.4)	94 (4.5)		92 (4.5)	94 (4.6)		
2008-2009	500 (11.2)	226 (10.7)		213 (10.3)	221 (10.7)		
2010-2011	811 (18.2)	421 (19.9)		396 (19.2)	409 (19.8)		
2012-2013	1086 (24.4)	529 (25.1)		517 (25.1)	516 (25.0)		
2014-2016	1598 (35.9)	814 (38.6)		814 (39.5)	794 (38.5)		
Age at diagnosis, years							
18–35	30 (0.7)	9 (0.4)	0.4	8 (0.4)	9 (0.4)	1.0	
36–50	412 (9.3)	203 (9.6)		201 (9.8)	194 (9.4)		
51–65	1942 (43.7)	961 (45.6)		939 (45.6)	935 (45.4)		
66–80	1898 (42.7)	864 (41.0)		834 (40.5)	851 (41.3)		
80+	163 (3.7)	72 (3.4)		79 (3.8)	72 (3.5)		
Sex							
Male	2236 (50.3)	1102 (52.2)	0.1	1073 (52.1)	1079 (52.4)	0.9	
Female	2213 (49.7)	1009 (47.8)		988 (47.9)	982 (47.6)		
Race							
White	3819 (85.8)	1905 (90.2)	< 0.001	1860 (90.2)	1857 (90.1)	1.0	
Black	447 (10.0)	138 (6.5)		133 (6.5)	136 (6.6)		
Other	183 (4.1)	68 (3.2)		68 (3.3)	68 (3.3)		
CDCC score		• •			• •		
0–1	4136 (93.0)	1973 (93.5)	0.5	1919 (93.1)	1924 (93.4)	0.8	
2+	313 (7.0)	138 (6.5)		142 (6.9)	137 (6.6)		

TABLE 1 (continued)

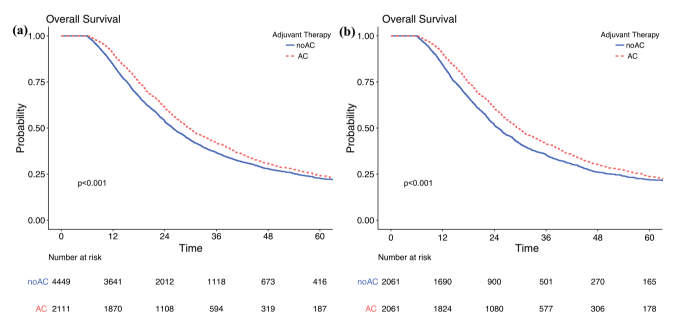
	Unmatched cohort			Matched cohort		
	noAC [n = 4449]	AC $[n = 2111]$	p value	noAC [n = 2061]	AC $[n = 2061]$	p value
Insurance status						
Uninsured	87 (2.0)	32 (1.5)	< 0.001	32 (1.6)	31 (1.5)	0.8
Private insurance	1915 (43.0)	1029 (48.7)		964 (46.8)	997 (48.4)	
Medicaid	214 (4.8)	81 (3.8)		84 (4.1)	80 (3.9)	
Medicare	2022 (45.4)	924 (43.8)		929 (45.1)	910 (44.2)	
Other/unknown	211 (4.7)	45 (2.1)		52 (2.5)	43 (2.1)	
Education level, %						
> 21	570 (12.8)	233 (11.0)	0.012	220 (10.7)	226 (11.0)	0.8
13–20.9	1063 (23.9)	472 (22.4)		484 (23.5)	463 (22.5)	
7–12.9	1563 (35.1)	739 (35.0)		732 (35.5)	726 (35.2)	
< 7	1253 (28.2)	667 (31.6)		625 (30.3)	646 (31.3)	
Medical income, US%						
≤ \$47,999	1704 (38.3)	712 (33.7)	0.001	707 (34.3)	699 (33.9)	0.8
\$48,000-\$62,999	1223 (27.5)	596 (28.2)		587 (28.5)	577 (28.0)	
\$63,000+	1522 (34.2)	803 (38.0)		767 (37.2)	785 (38.1)	
Residence						
Metro	3572 (80.3)	1678 (79.5)	0.9	1628 (79.0)	1640 (79.6)	1.0
Urban	683 (15.4)	333 (15.8)		331 (16.1)	324 (15.7)	
Rural	71 (1.6)	37 (1.8)		39 (1.9)	37 (1.8)	
Unknown	123 (2.8)	63 (3.0)		63 (3.1)	60 (2.9)	
Adjuvant chemotherapy						
Single agent	1185 (26.6)	628 (29.7)	< 0.001	603 (29.3)	611 (29.6)	1.0
Multi agent	2741 (61.6)	1385 (65.6)		1359 (65.9)	1352 (65.6)	
Unknown	523 (11.8)	98 (4.6)		99 (4.8)	98 (4.8)	
Neoadjuvant radiotherapy						
None	1540 (34.6)	845 (40.0)	< 0.001	846 (41.0)	828 (40.2)	0.6
NART	2909 (65.4)	1266 (60.0)		1215 (59.0)	1233 (59.8)	
Type of surgery						
Distal pancreatectomy	3317 (74.6)	1667 (79.0)	< 0.001	1596 (77.4)	1621 (78.7)	0.4
Pancreaticoduodenectomy	1132 (25.4)	444 (21.0)		465 (22.6)	440 (21.3)	
Tumor grade	, ,	,		, ,	` ,	
Well	397 (8.9)	150 (7.1)	0.006	156 (7.6)	147 (7.1)	1.0
Moderate	1389 (31.2)	724 (34.3)		708 (34.4)	712 (34.5)	
Poor	874 (19.6)	435 (20.6)		424 (20.6)	429 (20.8)	
Anaplastic	1789 (40.2)	802 (38.0)		773 (37.5)	773 (37.5)	
AJCC pathological T stage	, ,	,		, ,	` ,	
TO	1066 (24.0)	308 (14.6)	< 0.001	289 (14.0)	278 (13.5)	1.0
T1	490 (11.0)	233 (11.0)		223 (10.8)	229 (11.1)	
T2	483 (10.9)	258 (12.2)		247 (12.0)	256 (12.4)	
T3	2288 (51.4)	1246 (59.0)		1241 (60.2)	1233 (59.8)	
T4	122 (2.7)	66 (3.1)		61 (3.0)	65 (3.2)	
AJCC pathological N stage	122 (217)	00 (8.1)		01 (5.0)	00 (0.2)	
N0	2866 (64.4)	1088 (51.5)	< 0.001	1057 (51.3)	1056 (51.2)	1.0
N1	1148 (25.8)	720 (34.1)	\ 0.001	720 (34.9)	711 (34.5)	1.0
N2	301 (6.8)	189 (9.0)		177 (8.6)	186 (9.0)	
N3	134 (3.0)	114 (5.4)		107 (5.2)	108 (5.2)	
Margin status	157 (5.0)	117 (3.7)		107 (3.2)	100 (3.2)	
mangin sunus	3797 (85.3)	1772 (83.9)	0.1	1732 (84.0)	1729 (83.9)	0.9

Table 1 (continued)

	Unmatched cohort		Matched cohort			
	noAC [n = 4449]	AC [ <i>n</i> = 2111]	p value	noAC [n = 2061]	AC [ <i>n</i> = 2061]	p value
Positive	652 (14.7)	339 (16.1)		329 (16.0)	332 (16.1)	
Lymphovascular invasion						
Absent	3608 (81.1)	1581 (74.9)	< 0.001	1532 (74.3)	1538 (74.6)	0.9
Present	841 (18.9)	530 (25.1)		529 (25.7)	523 (25.4)	
Length of stay						
Median (IQR)	9.0 (11.0)	8.0 (8.0)	< 0.001	9.0 (9.0)	8.0 (8.0)	0.8

Data are expressed as n (%)

AC adjuvant chemotherapy, AJCC American Joint Committee on Cancer, CDCC Charlson–Deyo comorbidity, IQR interquartile range, NART neoadjuvant radiotherapy, noAC no adjuvant chemotherapy



**FIG. 1.** Overall survival of adjuvant chemotherapy following resection for pancreatic ductal adenocarcinoma in (a) unmatched and (b) matched cohorts. *AC* adjuvant chemotherapy, *noAC* no adjuvant chemotherapy

Association of Adjuvant Chemotherapy and Neoadjuvant Radiotherapy with Survival

Additional analyses were performed to further understand the impact of AC in patients by surgery type (i.e. PD or DP). In unadjusted analysis, there were significant differences in survival between AC and noAC patients in those who underwent PD (median 27.4 vs. 23.9 months; p = 0.002) and those who underwent DP (median 30.2 vs. 25.3 months; p < 0.001). On multivariable analyses modeling the interaction between receipt of AC and radiotherapy, a survival benefit was again seen for patients who underwent PD (HR 0.77, 95% CI 0.66–0.91; p < 0.001) and DP (HR 0.82, 95% CI 0.75–0.90; p < 0.001) (Table 4). As a sensitivity analysis, we

performed two separate multivariable analyses in cohorts including only those who underwent PD and DP, respectively. These analyses confirmed the same findings (Table 2).

#### DISCUSSION

Although current national guidelines from ASCO and NCCN recommend considering additional AC in patients following NAT and surgery, no high-level data exist to support this recommendation. <sup>4,5</sup> This national population-based cohort utilizing the NCDB and including 6560 patients who received NAC followed by resection for PDAC, demonstrated AC was associated with improved overall survival after accounting for potential biases

TABLE 2 Association of adjuvant chemotherapy with overall survival of patients with resected pancreatic ductal adenocarcinoma in unmatched and matched cohorts and stratified by nodal status and margin status for matched cohorts from a multivariable Cox regression model

Cohort	Chemotherapy	Median survival (IQR), months	HR (95% CI)	<i>p</i> -Value
All patients				
Unmatched	noAC	25.9 (25.1–26.7)	Ref	< 0.001
	AC	29.5 (28.5–30.9)	0.82 (0.76–0.87)	
Matched	noAC	24.9 (23.9–26.0)	Ref	< 0.001
	AC	29.4 (28.4–30.8)	0.81 (0.75–0.88)	
Stratified by nodal status in the matched cohort				
N0	noAC	29.4 (27.6–31.4)	Ref	< 0.001
	AC	34.0 (31.4–37.7)	0.80 (0.72–0.90)	
N1	noAC	22.1 (20.8–23.5)	Ref	< 0.001
	AC	28.2 (26.4–30.2)	0.76 (0.67–0.86)	
N2/3	noAC	20.0 (18.5–22.7)	Ref	0.9
	AC	21.9 (19.8–24.1)	1.01 (0.83–1.24)	
Stratified by margin status in the matched cohort				
R0	noAC	26.7 (25.6–28.7)	Ref	< 0.001
	AC	31.2 (29.8–34.0)	0.82 (0.75–0.89)	
R1	noAC	18.1 (16.6–19.8)	Ref	0.007
	AC	22.1 (19.6–24.0)	0.77 (0.64–0.93)	
Stratified by neoadjuvant radiotherapy status in the matched cohort				
No neoadjuvant radiotherapy	noAC	24.7 (22.8–27.5)	Ref	0.009
	AC	31.4 (28.6–35.5)	0.84 (0.74–0.96)	
Neoadjuvant radiotherapy	noAC	24.9 (23.8–26.2)	Ref	< 0.001
	AC	28.9 (27.4–30.5)	0.80 (0.73–0.88)	
Stratified by surgery type				
Pancreaticoduodenectomy	noAC	23.9 (21.5–25.7)	Ref	0.002
	AC	27.4 (25.6–30.2)	0.77 (0.66–0.91)	
Distal pancreatectomy	noAC	25.3 (24.0–26.7)	Ref	< 0.001
	AC	30.2 (28.6–31.9)	0.82 (0.75–0.89)	

AC adjuvant chemotherapy, CI confidence interval, HR hazard ratio, IQR interquartile range, noAC no adjuvant chemotherapy, Ref referent

through PSM. Stratified analyses by nodal and margin status demonstrated survival benefit of AC remained, even in patients considered to be at low risk of recurrence (i.e. margin-negative and node-negative disease. Furthermore, stratified analysis by receipt of NART also demonstrated a similar protective effect of AC. These findings collectively

suggest that completing planned systemic therapy should be considered after NAC and surgery for PDAC, whenever possible.

To date, evidence on whether AC after NAT and surgery confers incremental advantage remains an ongoing debate as several retrospective studies on this matter have yielded conflicting results. <sup>13–18</sup> Previous NCDB analyses by

**TABLE 3** Multivariable Cox regression model of survival of patients with resected pancreatic ductal adenocarcinoma in the matched cohort

	HR (95% CI)	<i>p</i> -Value
Center volume		
Quintile 1	Ref	
Quintile 2	0.99 (0.85-1.15)	0.9
Quintile 3	0.98 (0.84-1.16)	0.8
Quintile 4	0.94 (0.80-1.11)	0.5
Quintile 5	0.78 (0.65-0.92)	0.004
Facility type		
Community	Ref	
Academic	1.05 (0.93-1.19)	0.4
Others	0.99 (0.85-1.17)	0.9
Facility location		
Northeast	Ref	
South	1.34 (1.20–1.49)	< 0.001
Midwest	1.18 (1.06–1.32)	0.003
West	1.13 (0.99–1.29)	0.1
Unknown	0.72 (0.31–1.64)	0.4
Hospital distance, miles		
< 12.5	Ref	
12.5-49.9	1.06 (0.97–1.17)	0.2
≥ 50	1.05 (0.93–1.18)	0.4
Year of diagnosis	,	
2004–2005	Ref	
2006–2007	0.69 (0.50-0.95)	0.025
2008-2009	0.83 (0.62–1.12)	0.2
2010-2011	0.72 (0.53–0.96)	0.027
2012–2013	0.62 (0.46–0.83)	0.001
2014–2016	0.76 (0.56–1.02)	0.1
Age at diagnosis, years	,	
18–35	Ref	
36–50	0.55 (0.20–1.56)	0.3
51–65	0.59 (0.21–1.70)	0.3
66–80	0.60 (0.21–1.72)	0.3
80+	0.72 (0.25–2.10)	0.5
Sex	,	
Male	Ref	
Female	0.97 (0.89–1.04)	0.4
Race	,	
White	Ref	
Black	0.90 (0.76–1.06)	0.2
Other	0.98 (0.78–1.22)	0.9
CDCC score	· · · · · · · · · · · · · · · · · · ·	
0–1	Ref	
2+	1.08 (0.93–1.25)	0.3
Insurance status	(0.20 1.20)	3.5
Uninsured	Ref	
Private insurance	0.80 (0.59–1.09)	0.2
Medicaid	0.77 (0.54–1.10)	0.2

TABLE 3 (continued)

	HR (95% CI)	<i>p</i> -Value
Medicare	0.88 (0.64–1.21)	0.4
Unknown	0.83 (0.57–1.22)	0.4
Education level, %		
> 21%	Ref	
13–20.9	1.09 (0.94–1.26)	0.3
7–12.9	1.17 (1.01–1.36)	0.034
< 7	1.22 (1.03–1.44)	0.021
Median income, US%		
≤ \$47,999	Ref	
\$48,000–\$62,999	0.89 (0.80-0.99)	0.037
\$63,000+	0.85 (0.75–0.97)	0.012
Residence		
Metro	Ref	
Urban	0.94 (0.84–1.07)	0.4
Rural	0.73 (0.53–1.01)	0.1
Unknown	1.01 (0.80–1.27)	0.9
Neoadjuvant chemotherapy agent		
Single agent	Ref	
Multi-agent	0.85 (0.78-0.93)	< 0.001
Unknown	0.89 (0.74–1.07)	0.2
Neoadjuvant radiotherapy		
No	Ref	
Yes	1.06 (0.98–1.16)	0.2
Type of surgery		
Distal pancreatectomy	Ref	
Pancreaticoduodenectomy	1.07 (0.97–1.17)	0.2
Tumor grade		
Well	Ref	
Moderate	1.54 (1.31–1.82)	< 0.001
Poor	1.89 (1.59–2.25)	< 0.001
Anaplastic	1.43 (1.21–1.69)	< 0.001
AJCC pathological T stage		
T0	Ref	
T1	0.88 (0.74–1.05)	0.2
T2	0.92 (0.78-1.08)	0.3
T3	1.10 (0.97–1.25)	0.1
T4	1.29 (1.02–1.63)	0.036
AJCC pathological N stage		
N0	Ref	
N1	1.21 (1.10-1.32)	< 0.001
N2	1.53 (1.32–1.76)	< 0.001
N3	1.66 (1.39–1.98)	< 0.001
Margin status		
Negative	Ref	
Positive	1.56 (1.42–1.73)	< 0.001
Lymphovascular invasion		
Absent	Ref	
Present	1.12 (1.01–1.24)	0.028

Table 3 (continued)

	HR (95% CI)	p-Value	
Length of stay			
Mean (SD)	1.01 (1.00–1.01)	0.003	
Adjuvant therapy			
None	Ref		
Yes	0.81 (0.75–0.88)	< 0.001	

AJCC American Joint Committee on Cancer, CDCC Charlson–Deyo comorbidity, CI confidence interval, HR hazard ratio, Ref referent, SD standard deviation

Swords et al. 13 and de Geus et al. 14 have attempted to study the benefit of AC following NAT. Swords et al. 13 reported that AC was associated with improved survival only in patients with a lymph node ratio between 0.01 and 0.14, not in patients with node-negative disease or a lymph node ratio > 0.15. On the other hand, de Geus et al. <sup>14</sup> found AC was not associated with improved survival in comparable settings, even in patients with node-positive or marginpositive disease. However, both studies had limitations, which were addressed in this study. First, our study utilized a contemporary edition of NCDB, which allowed for a larger cohort to be scrutinized, particularly as NAT use increased over time. For example, in the study by de Geus et al. 14 833 noAC and 524 AC patients were included, compared with 4449 and 2111, respectively, in this current study. This likely increased power and generalizability. Second, the present study adopted PSM to adjust for selection biases for receiving either AC or noAC to create well-balanced cohorts. Finally, previous analyses 13,14 did not establish any survival benefit for AC in patients with low-risk disease, which was demonstrated in the present study.

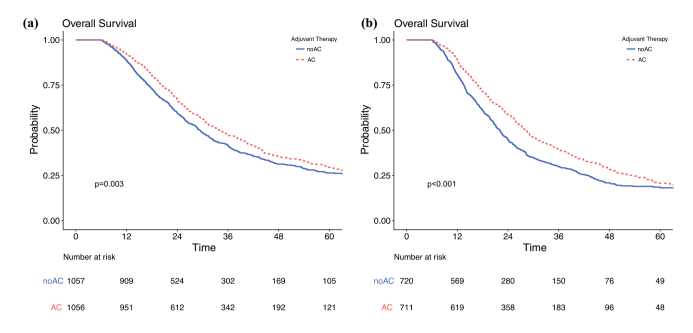
Previous single-center institutional series on this topic have also been limited by very small sample sizes, precluding valid conclusions to be drawn. <sup>15–18</sup> A recent retrospective analyses by Perri et al. that included 245 patients demonstrated noAC patients had received more chemotherapy cycles preoperatively (median 4 vs. 3), had larger tumors (2.8 vs. 2.4 cm), and had higher preoperative cancer antigen (CA) 19-9 levels (32 vs. 24 U/mL). <sup>18</sup> After PSM, AC was found to be associated with improved recurrence-free survival (median 17 vs. 12 months) but not

overall survival (median 42 vs. 32 months). Although the authors concluded that AC should be universally considered after NAT and surgery, subgroup analyses by nodal or margin status are lacking. In a previous study from the same group, AC was associated with improved survival only in patients with a low lymph ratio and in node-negative disease. <sup>15</sup>

The presence of high-risk factors, such as nodal involvement or positive margins, is commonly used to select patients for adjuvant therapy, as evidence by the distribution of AC use in the unmatched cohort. However, systemic and local recurrence in patients perceived to have low-risk disease are still as high as  $40\%^{26,27}$  and 50%. respectively. 28,29 This suggests that clinicians may underestimate the risk of relapse in PDAC patients perceived to have favorable pathologic features after resection. 26,27 To the authors' knowledge, no published studies have explored the role of AC specifically in patients with margin-negative or node-negative disease. These data suggest that AC is likely beneficial in most patients and is consistent with the notion that localized PDAC should be approached as a systemic disease.<sup>30</sup> While no specific criteria exist to aid in identifying patients most likely to benefit from AC, a recent study by Liu et al. found that CA19-9 response to NAT predicted incremental survival advantage, with patients only garnering benefit when levels failed to drop below 50% from baseline.<sup>31</sup> While these results are important to bear in mind, this study supports considering AC universally in the absence of higher-quality data.

This study has several key strengths that should be emphasized. First, the study utilizes a large and contemporary national cohort that probably resembles real-world trends and outcomes. Indeed, NCDB is particularly suited to test this study's stated hypothesis given an impressive level of granularity and high-quality standards. Second, detailed subgroup analyses in patients with negative lymph nodes, negative resection margins, and those who received NART were conducted to better inform decision making in clinically relevant scenarios. Lastly, potential treatment selection bias is minimized through PSM and subgroup and sensitivity analyses. We believe this lends further credibility to the stated conclusions.

Limitations of this analysis should also be acknowledged. First, this was a non-randomized, retrospective cohort study that is susceptible to bias. We attempted to minimize bias through PSM but potential unmeasured covariates may have contributed to the observed outcomes.



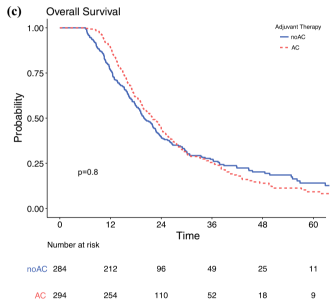


FIG. 2. Overall survival of adjuvant chemotherapy following resection for pancreatic ductal adenocarcinoma stratified by nodal status in matched cohorts (a) N0, (b) N1, and (c) N2/3. AC adjuvant chemotherapy, noAC no adjuvant chemotherapy

Furthermore, patients with survival of < 6 months were excluded, primarily to account for patients who did not survive long enough to receive AC; however, it is possible that doing so also excluded patients who died due to AC-related complications, although this is likely to be a small group. Second, 60% of patients in this analysis received NART in addition to NAC. This may therefore limit generalizability as NART is uncommonly used in resectable tumors. To accommodate for that, we conducted a separate analysis according to receipt of NART and confirmed the advantage of AC in this subgroup. Third, details on AC agent(s) were limited to single- or multi-

agent categories, with no further information. Nevertheless, a majority of patients in this study received multi-agent AC, which perhaps indicates that most NAC patients ultimately recover well from pancreatectomy and are eligible to receive more than single-agent regimens. Finally, it is important to note that the number of NAC cycles received is unclear from NCDB and that a recommendation for additional AC likely does not apply to patients who received longer courses of NAC, as endorsed by some investigators.<sup>32</sup> Hence, this study cannot provide specific guidance with respect to the duration of either preoperative or postoperative therapy, differentiation between

TABLE 4 Multivariable Cox regression model of survival of patients with resected pancreatic ductal adenocarcinoma in the matched cohort, with interactions between chemotherapy and nodal status and margin status

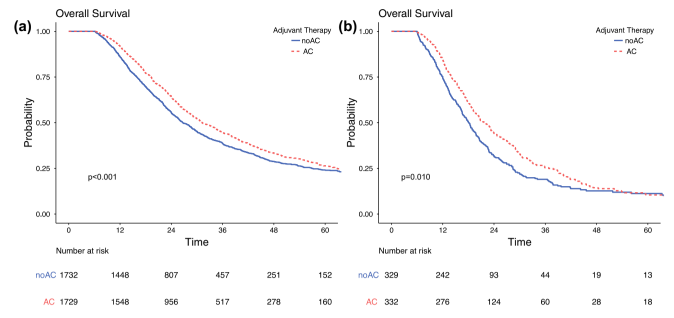
	HR (95% CI)	p Value
Interaction by nodal status		
Adjuvant chemotherapy * AJCC pathological N stage		
N0 + noAC	Ref	0.001
R0 + noAC	0.81 (0.68-0.97)	
R0 + AC	1.26 (1.12–1.42)	
R1 + noAC	0.76 (0.67-0.86)	
R1 + AC	1.56 (1.22–2.00)	
N2/3 + AC	0.99 (0.82-1.14)	
Interaction by margin status		
Adjuvant chemotherapy * margin status		
R0 + noAC	Ref	< 0.001
R0 + AC	0.83 (0.76-0.90)	
R1 + noAC	1.62 (1.41–1.86)	
R1 + AC	0.73 (0.57-0.93)	
Interaction by neoadjuvant radiotherapy		
Adjuvant chemotherapy * neoadjuvant radiotherapy		
noNART + noAC	Ref	< 0.001
noNART + AC	0.85 (0.75-0.96)	
NART + noAC	1.10 (0.98-1.23)	
NART + AC	0.74 (0.60-0.90)	
Interaction by surgery type		
Adjuvant chemotherapy * neoadjuvant radiotherapy		
DP + noAC	Ref	< 0.001
DP + AC	0.82 (0.75-0.90)	
PD + noAC	1.09 (0.96-1.23)	
PD + AC	0.77 (0.66-0.91)	

AC adjuvant chemotherapy, AJCC American Joint Committee on Cancer, CI confidence interval, DP distal pancreatectomy, HR hazard ratio, NART neoadjuvant radiotherapy, noNART no neoadjuvant radiotherapy, noAC no adjuvant chemotherapy, PD pancreaticoduodenectomy, Ref referent

chemotherapy from chemoradiation therapy versus chemotherapy plus chemoradiation, the subsets of patients for whom these treatments are most beneficial, and the extent to which histopathologic analysis of the surgical specimen may inform the postoperative treatment regimen. Ongoing randomized trials will likely answer these questions.

#### **CONCLUSION**

Adjuvant chemotherapy following resection is associated with improved long-term survival in patients receiving NAC, even in margin-negative and node-negative disease. These findings suggest that AC should be considered in patients who did not complete all intended systemic treatments upfront, whenever permissible.



**FIG. 3.** Overall survival of adjuvant chemotherapy following resection for pancreatic ductal adenocarcinoma stratified by margin status in matched cohorts (a) R0 and (b) R1. AC adjuvant chemotherapy, noAC no adjuvant chemotherapy

**FUNDING** No funding was received for this study.

**OPEN ACCESS** This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

#### REFERENCES

- Neoptolemos JP, Palmer DH, Ghaneh P, et al. Comparison of adjuvant gemcitabine and capecitabine with gemcitabine monotherapy in patients with resected pancreatic cancer (ESPAC-4): a multicentre, open-label, randomised, phase 3 trial. *Lancet*. 2017;389(10073):1011–24. https://doi.org/10.1016/S014 0-6736(16)32409-6.
- Conroy T, Hammel P, Hebbar M, et al. FOLFIRINOX or gemcitabine as adjuvant therapy for pancreatic cancer. N Engl J Med. 2018;379(25):2395–406. https://doi.org/10.1056/NEJMoa 1809775.
- Murphy JE, Wo JY, Ryan DP, et al. Total neoadjuvant therapy with FOLFIRINOX followed by individualized chemoradiotherapy for borderline resectable pancreatic adenocarcinoma: a phase 2 clinical trial. *JAMA Oncol*. 2018;4(7):963–9. https://doi.org/10. 1001/jamaoncol.2018.0329.
- Tempero MA, Malafa MP, Al-Hawary M, et al. Pancreatic adenocarcinoma, version 2.2017, NCCN clinical practice guidelines in oncology. *J Natl Compr Cancer Netw.* 2017;15(8):1028–61. h ttps://doi.org/10.6004/jnccn.2017.0131.

- Khorana AA, Mangu PB, Berlin J, et al. Potentially curable pancreatic cancer: American society of clinical oncology clinical practice guideline update. *J Clin Oncol*. 2017;35(20):2324–8. h ttps://doi.org/10.1200/JCO.2017.72.4948.
- Janssen QP, Buettner S, Suker M, et al. Neoadjuvant FOLFIR-INOX in patients with borderline resectable pancreatic cancer: a systematic review and patient-level meta-analysis. *J Natl Cancer Inst*. 2019;111(8):782–94. https://doi.org/10.1093/jnci/djz073.
- da Costa Jr WL, Tran Cao HS, Massarweh NN. Neoadjuvant treatment for patients with localized pancreatic adenocarcinoma: Are we there yet? *JAMA Oncol*. 2020;6(8):1163–4. https://doi. org/10.1001/jamaoncol.2020.0562.
- Oettle H, Neuhaus P, Hochhaus A, et al. Adjuvant chemotherapy with gemcitabine and long-term outcomes among patients with resected pancreatic cancer: the CONKO-001 randomized trial. *JAMA*. 2013;310(14):1473–81. https://doi.org/10.1001/jama.201 3.279201.
- Tzeng CW, Tran Cao HS, Lee JE, et al. Treatment sequencing for resectable pancreatic cancer: influence of early metastases and surgical complications on multimodality therapy completion and survival. *J Gastrointest Surg*. 2014;18(1):16–24. https://doi.org/ 10.1007/s11605-013-2412-1 (discussion 24-5).
- Versteijne E, Suker M, Groothuis K, et al. Preoperative chemoradiotherapy versus immediate surgery for resectable and borderline resectable pancreatic cancer: results of the dutch randomized phase III PREOPANC trial. *J Clin Oncol*. 2020;38(16):1763–73. https://doi.org/10.1200/JCO.19.02274.
- Katz MHG, Ou FS, Herman JM, et al. Alliance for clinical trials in oncology (ALLIANCE) trial A021501: preoperative extended chemotherapy vs. chemotherapy plus hypofractionated radiation therapy for borderline resectable adenocarcinoma of the head of the pancreas. *BMC Cancer*. 2017;17(1):505. https://doi.org/10. 1186/s12885-017-3441-z.
- Katz MH, Shi Q, Ahmad SA, et al. Preoperative modified FOLFIRINOX treatment followed by capecitabine-based chemoradiation for borderline resectable pancreatic cancer: alliance for clinical trials in oncology trial A021101. *JAMA Surg*. 2016;151(8):e161137. https://doi.org/10.1001/jamasurg.2016. 1137.

- Swords DS, Francis SR, Lloyd S, et al. Lymph node ratio in pancreatic adenocarcinoma after preoperative chemotherapy vs. preoperative chemoradiation and its utility in decisions about postoperative chemotherapy. *J Gastrointest Surg*. 2019;23(7):1401–13. https://doi.org/10.1007/s11605-018-3953-0.
- de Geus SWL, Kasumova GG, Eskander MF, et al. Is neoadjuvant therapy sufficient in resected pancreatic cancer patients? A national study. *J Gastrointest Surg*. 2018;22(2):214–25. https://doi.org/10.1007/s11605-017-3541-8.
- Roland CL, Katz MH, Tzeng CW, et al. The addition of postoperative chemotherapy is associated with improved survival in patients with pancreatic cancer treated with preoperative therapy. *Ann Surg Oncol*. 2015;22(Suppl 3):S1221–8. https://doi.org/10. 1245/s10434-015-4854-z.
- Barnes CA, Krepline AN, Aldakkak M, et al. Is adjuvant therapy necessary for all patients with localized pancreatic cancer who have received neoadjuvant therapy? *J Gastrointest Surg*. 2017;21(11):1793–803. https://doi.org/10.1007/s11605-017-3544
- Groot VP, Blair AB, Gemenetzis G, et al. Recurrence after neoadjuvant therapy and resection of borderline resectable and locally advanced pancreatic cancer. Eur J Surg Oncol. 2019;45(9):1674–83. https://doi.org/10.1016/j.ejso.2019.04.007.
- Perri G, Prakash L, Qiao W, et al. Postoperative chemotherapy benefits patients who received preoperative therapy and pancreatectomy for pancreatic adenocarcinoma. *Ann Surg*. 2020;271(6):996–1002. https://doi.org/10.1097/SLA. 0000000000003763.
- van Roessel S, van Veldhuisen E, Klompmaker S, et al. Evaluation of adjuvant chemotherapy in patients with resected pancreatic cancer after neoadjuvant FOLFIRINOX treatment. *JAMA Oncol.* 2020;6(11):1–8. https://doi.org/10.1001/jamaoncol. 2020.3537.
- Bilimoria KY, Bentrem DJ, Ko CY, et al. Validation of the 6th edition AJCC pancreatic cancer staging system: report from the national cancer database. *Cancer*. 2007;110(4):738–44. https://d oi.org/10.1002/cncr.22852.
- Merkow RP, Rademaker AW, Bilimoria KY. Practical guide to surgical data sets: national cancer database (NCDB). *JAMA Surg*. 2018;153(9):850–1. https://doi.org/10.1001/jamasurg.2018.0492.
- Giobbie-Hurder A, Gelber RD, Regan MM. Challenges of guarantee-time bias. *J Clin Oncol*. 2013;31(23):2963–9. https://doi.org/10.1200/JCO.2013.49.5283.
- Austin PC. The use of propensity score methods with survival or time-to-event outcomes: reporting measures of effect similar to those used in randomized experiments. Stat Med. 2014;33(7):1242–58. https://doi.org/10.1002/sim.5984.
- 24. Kamarajah SK, Sonnenday CJ, Cho CS, et al. Association of adjuvant radiotherapy with survival after margin-negative

- resection of pancreatic ductal adenocarcinoma: a propensity-matched national cancer database (NCDB) analysis. *Ann Surg.* 2021;273(3):587–94. https://doi.org/10.1097/SLA. 00000000000003242.
- 25. Kamarajah SK, Phillips AW, Hanna GB, Low DE, Markar SR. Definitive chemoradiotherapy compared to neoadjuvant chemoradiotherapy with esophagectomy for locoregional esophageal cancer: national population-based cohort study. *Ann Surg.* 2019. https://doi.org/10.1097/SLA.00000000000003941 (Epub 19 May 202).
- Kim RD, Kundhal PS, McGilvray ID, et al. Predictors of failure after pancreaticoduodenectomy for ampullary carcinoma. *J Am Coll Surg.* 2006;202(1):112–9. https://doi.org/10.1016/j.jamcollsurg.2005.08.002.
- Lee JH, Whittington R, Williams NN, et al. Outcome of pancreaticoduodenectomy and impact of adjuvant therapy for ampullary carcinomas. *Int J Radiat Oncol Biol Phys.* 2000;47(4):945–53. https://doi.org/10.1016/s0360-3016(00)00537-x
- Junrungsee S, Kittivarakul E, Ko-iam W, Lapisatepun W, Sandhu T, Chotirosniramit A. Prognostic factors and survival of patients with carcinoma of the ampulla of vater after pancreaticoduodenectomy. *Asian Pac J Cancer Prev.* 2017;18(1):225–9. https://doi.org/10.22034/APJCP.2017.18.1.225.
- 29. de Castro SM, Kuhlmann KF, van Heek NT, et al. Recurrent disease after microscopically radical (R0) resection of periampullary adenocarcinoma in patients without adjuvant therapy. J Gastrointest Surg. 2004;8(7):775–84. https://doi.org/10.1016/j.gassur.2004.08.006 (discussion 784).
- Epelboym I, Zenati MS, Hamad A, et al. Analysis of perioperative chemotherapy in resected pancreatic cancer: identifying the number and sequence of chemotherapy cycles needed to optimize survival. *Ann Surg Oncol.* 2017;24(9):2744–51. https://doi.org/10.1245/s10434-017-5975-3.
- Liu H, Zenati MS, Rieser CJ, et al. CA19-9 change during neoadjuvant therapy may guide the need for additional adjuvant therapy following resected pancreatic cancer. *Ann Surg Oncol*. 2020;27(10):3950–60. https://doi.org/10.1245/s10434-020-08468
- Truty MJ, Kendrick ML, Nagorney DM, et al. Factors predicting response, perioperative outcomes, and survival following total neoadjuvant therapy for borderline/locally advanced pancreatic cancer. *Ann Surg.* 2021;273(2):341–9. https://doi.org/10.1097/S LA.00000000000003284.

**Publisher's Note** Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.