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Endoscopic Ultrasound-Guided Radiofrequency Ablation as an Future Alternative to Pancreatectomy for Pancreatic Metastases from Renal Cell Carcinoma: A Prospective Study

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Abstract: Background: Pancreatic metastases (PM) from renal cell carcinoma (mRCC) are rare, are associated with a long survival. Focal treatment in order to control oligo-metastatic disease and avoid systemic therapy is a standard in RCC. However, pancreatic radiofrequency ablation remains a marginal and under evaluated technic. Standard treatment remains pancreatectomy with hazardous outcomes. We report here the largest series of endoscopic radiofrequency ablation procedure then followed by CT-scan. We prospectively included 12 patients, median age was 70.5 years old and median size and number of PM at inclusion was 17 mm and 1 respectively. All 26 procedures performed went as planned and no immediate complications were experienced. After 27.7 months of follow-up, the 6- and 12-month focal control rates were 84% and 73% respectively. Two severe complications occurred after EUS-RFA but were totally resolved after specific treatment. In total, EUS-RFA is feasible and displayed an excellent focal control and remained pancreatectomy with hazardous outcomes. We report here the largest series of endoscopic radiofrequency ablation (EUS-RFA) on PM for patient treated at Institut Paoli-Calmettes for a mRCC. Patients with progressive PM was not an available guideline, and 73% respectively. Two severe complications occurred after EUS-RFA but were totally resolved after specific treatment. In total, EUS-RFA is feasible and displayed an excellent focal control and should be integrated in the arsenal to treat PM from mRCC in order to spare systemic therapy and/or pancreatic surgery.

Simple Summary: Glandular metastases and more precisely pancreatic metastases (PM) from renal cell carcinoma (mRCC) are associated with a long survival. Focal treatment in order to control oligo-metastatic disease and avoid systemic therapy is a standard in RCC. However, pancreatic radiofrequency ablation remains a marginal and under evaluated technic. Standard treatment remains pancreatectomy with hazardous outcomes. We report here the largest series of endoscopic radiofrequency ablation (EUS-RFA) on PM for patient treated at Institut Paoli-Calmettes for a mRCC. Patients with progressive PM was treated under general anesthesia with an ultrasound guided endoscopic radiofrequency ablation procedure then followed by CT-scan. We prospectively included 12 patients, median age was 70.5 years old and median size and number of PM at inclusion was 17 mm and 1 respectively. All 26 procedures performed went as planned and no immediate complications were experienced. After 27.7 months of follow-up, the 6- and 12-month focal control rates were 84% and 73% respectively. Two severe complications occurred after EUS-RFA but were totally resolved after specific treatment. In total, EUS-RFA is feasible and displayed an excellent focal control and should be integrated in the arsenal to treat PM from mRCC in order to spare systemic therapy and/or pancreatic surgery.

Keywords: Pancreatic metastases; Renal cell carcinoma; Endoscopic ultrasound-guided radiofrequency ablation; EUS-RFA; mRCC; Pancreatectomy; Focal therapy; Systemic therapy; VEGFR inhibitors; ECOG 0–1.

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For evaluable PM, the 6- and 12-month focal control rates were 84% and 73% respectively. One patient treated with TKI developed a paraduodenal abscess 2 months after EUS-RFA and another patient with biliary stent developed hepatic abscesses few days after EUS-RFA. No other severe side effects were experienced. Conclusions: in this series, which is the largest ever reported, we showed that EUS-RFA is feasible and yields an excellent local control rate for PM from mRCC. With manageable complications, it could be a valuable alternative to pancreatic surgery in well-selected patients.

**Keywords:** glandular metastases; endoscopy; radiofrequency ablation; renal cell carcinoma; pancreatic metastases

### 1. Introduction

Pancreatic metastases (PM) are rare, but when detected, renal cell carcinoma is the most frequent primary tumor [1]. PMs are detected in 1 to 6% of metastatic renal cell carcinoma (mRCC) and are frequently associated with other glandular metastases [2], which are known to confer a better outcome compared to liver, bones or brain metastases [3–5]. Although molecular mechanisms associated with long survival of patients presenting PM are poorly understood, primary tumors exhibit slow-growing characteristics (low Fuhrman/ISUP grade, long period between primary tumor and metastatic relapse) [1] or specific molecular/immune features (linear and monogenic primary to metastasis evolution, reduced effector T cell gene signatures and increased angiogenesis dependence) [6,7].

Focal therapies are now a standard treatment of oligometastic mRCC [8]. Pancreatic resection is the most common option to control isolated PM and to avoid compressive symptoms, whereas tyrosine kinase inhibitors (TKI) are preferentially used when other metastatic sites are detected [9,10].

However, these strategies present morbid side-effects and not all patients are fit enough to undergo pancreatectomy [1]. Pancreatic surgery for RCC with solitary PM has been extensively reported and 5-year survival rates range 50 to 75% [1,10–12]. However, this complex surgery has a post-operative mortality of around 2% and morbidity varies from 13.5 to 43% (pancreatic fistula, infections or diabetes) [9,13,14]. Thus, less invasive techniques need to be developed and validated.

Endoscopic Ultra-Sound (EUS)-guided radiofrequency ablation (RFA) is a microinvasive technique which need a classical therapeutic EUS scope (with 4 mm working channel) and a dedicated 19 Gauge EUS-RFA needle [15]. The EUS-RFA probe delivers locally a high-frequency (400–500 kHz) alternating (sinusoidal) current that generates electromagnetic high-temperature cell damages, limiting bleeding [16]. Percutaneous-RFA has shown efficacy to treat both primary and secondary liver tumors and recent reports showed successful transposition of this technique to pancreatic tumors [15,17]. The technique seems promising in hyper-vascularized tumors like RCC. However, large cohorts of RFA (external or endoscopic) in the treatment of PM are lacking and only case reports are available displaying heterogeneous results and toxicity profiles [18,19].

We report here the efficacy and safety of EUS-RFA in a series of patients with PM from RCC treated at the Institut Paoli-Calmettes (Comprehensive Cancer Center, Marseille, France).

### 2. Materials and Methods

This prospective, monocentric, non-controlled, single arm pilot study aimed to analyze the safety and the focal control of EUS-RFA used to treat PM from RCC.
2.1. Patients

Eligible patients was >18 years-old, had metastatic clear cell RCC with progressive but limited PM (<4 PM, maximum diameter: 25 mm) [17], ECOG 0–1, eligible to general anesthesia and treated or not for extra-PM disease. Extra PM metastases, when present, had to be limited and stable. The indications of EUS-RFA were validated during cancer multidisciplinary team meetings and all patients gave their written informed consent. The protocol was approved by the institutional review board of Institut Paoli-Calmettes (the IRB number of this study is: PRAM-1-IPC 2018-013, approval date: 4 July 2018).

2.2. EUS-RFA Procedure

EUS-RFA was performed under general anesthesia, with a linear EUS scope.

We used an EUS-RFA system (STARmed, Seoul, Korea) consisting of a 19-gauge needle electrodes covered with a sheath, except for the terminal 1 cm with a sharp conical tip for energy delivery, and an radiofrequency (RF) generator. This system features an internal cooling system which circulates a chilled saline solution through the needle electrode during the RFA procedure to prevent charring on the surface of the electrode and to improve the accuracy of the ablation. The lesion was punctured at 1 to 3 different points, and 50 W ablation power was delivered at the different sites under direct EUS control. Ablation time, related to tissue impedance measured in real time by the RF generator, was short (10–20 s but when PM was >20 mm, time was slightly longer from 30–60 s) (Supplement Video S1). Bioimpedance reflects the property of a tissue to conduct electricity. An impedance rise is an indirect sign of cellular disruption, tissue charring, and hampering of electrical conduction. When a sudden rise in impedance was registered, power delivery was stopped.

2.3. Prophylaxis and Evaluation of Immediate Complications

Before ablation, all patients received non-steroidal anti-inflammatory treatment (indomethacin by rectal route) to prevent acute pancreatitis and antibioprophylaxis using cefotaxime.

Patients were kept two days in the hospital for surveillance of pain, bleeding, perforation signs, fever or other acute pancreatitis symptoms.

Before general anesthesia, indomethacin and cefotaxime were administrated to prevent acute pancreatitis. EUS-RFA was performed with a linear EUS scope and a EUS-RFA system (STARmed, Seoul, Korea—19-gauge needle electrode). The lesion was punctured (at 1–3 points) (Supplement Video S1), ablation time was short (50 watt–30 s to 1 min) and controlled by tissue impedance measured in real time by the RF generator.

2.4. Tumor Response Assessment

CT-scan was performed 2 months after each EUS-RFA procedure, then every 3 to 6 months for follow-up. Response was assessed for each treated PM using size and contrast uptake (Figure 1). Complete response was defined by complete disappearance of the lesion or absence of any contrast uptake at the arterial phase [17]; partial response was a regression of 30% or more of the tumor contrast uptake, stable disease was defined with size change comprised between −30% and +20% of the contrast uptake of PM. Progressive disease was defined as a >20% increase in size of PM with contrast uptake or new compressive symptoms [20].
Figure 1. Following contrast enhancing PM with CT-scan before and after EUS-RFA. Abdomen CT-scan with contrast injection at artery phase, before (A) and after endoscopic radiofrequency ablation of PM in the body of the pancreas, displaying complete response without contrast enhancement in RFA scar (B) and at magnification (C). PM: pancreatic metastases, CT: computed tomography, EUS-RFA: Endoscopic Ultra-Sound guided RadioFrequency Ablation.

Focal control rate was defined as the sum of complete response, partial response and stable disease rates.

3. Results

Between June 2017 and December 2019, twelve patients underwent EUS-RFA for PM from mRCC, at the Institut Paoli-Calmettes (Marseille, France), in the Endoscopic and Interventional unit.

All had undergone past nephrectomy for RCC, 50% were female, and the median age at inclusion was 70.5 years-old (range 62–75). Median time between nephrectomy and PM was 13.6 years (range 0–22.2) (Table 1). For 58%, PM remained the unique metastatic site until the last evaluation. The mean size of PM was 17 mm (range 3–35) and 33% had an initial size >20 mm (range 22–35) (Supplement Table S1). One patient developed jaundice and required an endoscopic drain with a biliary stent.

Table 1. Baseline patients’ characteristics (n = 12).

| Baseline Patients’ Characteristics (n = 12) | n (%) |  |
|------------------------------------------|-------|---|
| Left Kidney                              | 6     | 60|
| Gender                                   |       |   |
| Female n (%)                             | 6     | 50|
| TNM                                      |       |   |
| T1/T2                                    | 9     | 75|
| T3/T4                                    | 1     | 8 |
| Unknown                                  | 2     | 17|
| Fuhrman Grade                            |       |   |
| I/II                                     | 7     | 58|
| III/IV                                   | 3     | 25|
| Unknown                                  | 2     | 17|
| IMDC                                     |       |   |
| Good/Intermediate                        | 4/5   | 33/42|
| Unknown                                  | 3     | 25|
| Systemic Treatment                       |       |   |
| Before EUS-RFA                           | VEGFR-TKI/ICI | 4/0 | 33|
| During EUS-RFA                           | VEGFR-TKI/ICI | 1/1 | 8 |
| After EUS-RFA                            | VEGFR-TKI/ICI | 3/0 | 25|
| No systemic treatment                    | 7     | 58|
Table 1. Cont.

| Baseline Patients’ Characteristics (n = 12) | n | % |
|-------------------------------------------|---|---|
| Other Metastatic Sites During RCC Evolution | None | 7 | 58 |
|                                            | Lung | 5 | 42 |
|                                            | Brain | 1 | 8 |
|                                            | Liver | 2 | 17 |
|                                            | Thyroid | 1 | 8 |
|                                            | Others | 2 | 17 |
| Median Time from Nephrectomy to 1st Metastasis | Years (range) | 11.6 (0–22.2) |
| Median Time from Nephrectomy to PM | Years (range) | 13.6 (0–22.2) |
| Pancreas as the 1st Metastatic Site | n (%) | 9 | 75 |
| Symptoms at PM Presentation | None | 9 | 75 |
|                               | Jaundice | 1 | 8 |
|                               | Pain | 2 | 17 |
| PM Localization at Presentation | Head | 7 | 33 |
|                               | Uncus | 4 | 19 |
|                               | Body | 6 | 29 |
|                               | Tail | 4 | 19 |
| Median PM SIZE at Presentation | Millimeters, (range) | 17 (3–35) |
| Median Number of PM at Presentation (range) | 1 | (1–4) |
| PM Number at Presentation (n, %) | 1 | 58 |
|                               | 2 | 17 |
|                               | 3–4 | 25 |

TNM: Tumor Node Metastases Classification, IMDC: International Metastatic RCC Database Consortium International, EUS-RFA: Endoscopic Ultra-Sound guided Radio Frequency Ablation, VEGFR-TKI: Vascular Endothelial Growth Factor Receptor-Tyrosine Kinase Inhibitor, ICI: Immune Check Point Inhibitor, RCC: Renal Cell Carcinoma, PM: pancreatic metastases.

Five patients (42%) had received a systemic TKI before the first EUS-RFA (Table 1). Three patients began a TKI new line after EUS-RFA, all for extra pancreatic progression. We performed 26 EUS-RFA sessions to treat a total of 21 PMs (Supplement Table S1). Nine patients underwent ≥2 EUS-RFA due to an incomplete necrosis of PM at first evaluation (Supplement Figure S1).

A second and third session for a same patient could be motivated by (1) the presence of untreated metastases in the first session (due to extremely low size or new lesion); (2) incomplete PM necrosis evaluated by CT-Scan defined by contrast enhanced remnant tissue.

3.1. Toxicties

All procedures achieved technical success with no immediate complication and all patients spent 2 days at hospital for each session.

Two patients experienced complications requiring hospitalization and intervention (grade IIIb Clavien-Dindo [21]). In the first patient, treated by TKI at the time of EUS-RFA, a duodenal abscess occurred two months after the 2nd EUS-RFA. In the second patient, the only one with a biliary stent, a symptomatic hepatic abscess occurred one week after the EUS-RFA. At the time of analysis, all adverse effects had been resolved, and no further complication was detected in any patient.

These two patients were the only ones in the cohort to continue TKI during EUS-RFA and to have a biliary stent. These observations had to be confirmed in a larger population, but patients treated with EUS-RFA should probably stop TKI and not have biliary stents before the procedure.
3.2. Focal Control and Survival

Median follow-up since the first EUS-RFA was 27.7 months (range 6.4–57.1). Among the 21 PMs treated, 40% had a complete response at 12 months. The 6-month and 12-month focal control rates were 84.2% and 73.3% respectively among evaluable PMs. Progressive disease was observed in only 15.8% and 26.7% of evaluable PMs at 6 and 12 months respectively (Table 2). All patients were alive at the time of analysis and median progression free survival for PM was 25.38 months (Figure 2).

Table 2. Objective response rates at 2, 6 and 12 for PM.

| Time Point (Evaluable Lesion) | CR (%) | PR (%) | SD (%) | PD (%) | Focal Control Rate (%) | N/E |
|------------------------------|--------|--------|--------|--------|------------------------|-----|
| 2-months (n = 21)            | 33.3   | 14.3   | 42.9   | 9.5    | 90.5                   | 0   |
| 6-months (n = 19)            | 26.3   | 31.6   | 26.3   | 15.8   | 84.2                   | 9.5 |
| 12-months (n = 15)           | 40     | 33.3   | 0      | 26.7   | 73.3                   | 28.6|

PM: pancreatic metastasis CR: complete response, PR: partial response, SD: stable disease PD: progression disease; N/E Not evaluable lesion (that have progressed or follow up is < to the evaluation time point); Focal Control Rate = CR + PR + SD.

Comparing PM > 20 mm to those <20 mm, the 6-month focal control rate was lower (92% vs. 71%) (Supplement Table S2) and the time to progression was shorter although the difference was not statistically significant (p = 0.06) (Supplement Figure S2). Tumor size was not significantly associated with a better response (OR 0.37, p = 0.43) in logistic regression analysis (Supplement Table S3).

After the last EUS-RFA, two patients received a new systemic therapy, one performed duodeno-pancreatectomy for PM (the same patient who developed paraduodenal abcess), and 4 patients underwent focal treatment of other RCC metastases (metastasectomy and/or RFA on the liver or lung). Focal directed therapies for oligometastatic disease are considered as a standard treatment as they improve both overall survival and cancer-specific survival in RCC [8,22,23]. Here, EUS-RFA was perfectly integrated in a multimodal focal therapy strategy, which is of high importance for this ‘slow growing tumor’ population.

4. Discussion

4.1. EUS-RFA for Managing Slow Growing Metastases and Avoiding Morbid Surgical Resection

In this study, among the 26 procedures performed, only two were associated with severe side effects in 2/12 patients (17%). Two teams published case reports of external RFA for RCC-PM with contrasted results: for Elias et al., the two patients treated with
RFA during laparotomy or laparoscopy to treat 2 to 3 PM ≤ 20 mm, complementary to the resection of other RCC metastases, both experienced severe pancreatitis leading to emergency surgery. The authors concluded that “in 2003, RFA is not yet usable and safe” [18]. However, for Carrafiello et al., a percutaneous RFA for a 20 mm PM was performed safely, resulting in no residual disease one year later [19].

In our series, no acute pancreatitis was observed with the systematic administration of indomethacin and cefotaxime in primary prevention.

Our data highlighted that EUS-RFA can achieve a good focal control of PM with only 15.8 and 26.7% of progression at 6 and 12 months. According to these results, EUS-RFA should be a key option to treat and control PM from RCC.

Finally, all the 4 patients that experienced PM progression after EUS-RFA underwent additional treatment including immune check-point inhibitors (ICI). Further data could also address whether EUS-RFA could trigger an antigenic reaction [16,24] and bypass the PM resistance mechanism to ICI [7].

The main limitation of our work is the low number of treated patients. However, with 26 procedures, this study provided relevant information about the good tolerance and performance of EUS-RFA for PM.

4.2. RFA and Immunotherapy

Focal treatment like radiation can trigger inflammatory cell activation, pictured in hepatocellular carcinoma (HCC), in which RFA is a standard treatment, and where studies observed a T cell response against tumor tissue and an increased number of systemic NK cells and cytotoxicity [25,26]. Recent immune monitoring in 80 patients with HCC < 5 cm reported that systemic variations of activated NK cells (NKp30+) after RFA was associated with tumor recurrence and with increased systemic CD8 Central Memory [24]. This overall activation of the immune system by RFA could be even more efficient in association with immune checkpoint inhibitors, which are now a standard of care for mRCC [27–29]. PM was recently proposed as a RCC metastatic localization associated with low response and survival with nivolumab [7]. Such an ablation with EUS-RFA could trigger an antigenic reaction and sensitize PMs to immune checkpoint inhibitors.

4.3. Study Limitations

Obviously, the low number of treated patients and the relatively short follow-up are the main limitations of our study, but with 26 procedures, this study provided relevant information about the good tolerance and performance of EUS-RFA for PM. Longer follow up will also provide more mature data, especially among tumors what are not evaluable (Table 2 and Supplement Table S2).

However, a larger comparative study might be hard to set-up because of the rarity of PM.

Furthermore, data from primary pancreatic tumors (adenocarcinomas or PNETS) reinforce the level of evidence of good tolerance and efficacy of this innovative procedure [17,30].

5. Conclusions

Even if further prospective data are needed to reinforce our findings, EUS-RFA to treat PM from RCC appears to be implemented as a promising option in the armamentarium of focal therapy in well selected patients.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/cancers13215267/s1, Figure S1: Study detail; Figure S2: Time until progression for all treated PM; Table S1: EUS-RFA procedures characteristics; Table S2: Focal control at 2, 6 and 12 months according to PM size; Table S3: Logistic regression of variables potentially associated with 6-months objective response; Video S1: Displays an EUS-RFA procedure for a 30 mm PM localized in the pancreas body from a 75 years old man. This was the second procedure for this patient, to treat a remaining hyper-vascular zone. The PM is presented as an heterogenous nodule with
hypo-echogenicity. The catheter is placed in front of the lesion before the puncture (time 0:05), the white arrow shows the tip of the RFA needle. RFA was performed inside the entire lesion, two times (time 0:25 to 0:38 and time 0:45 to 0:47) then the needle is removed. A contrast enhanced endoscopic-echosonography is performed after the procedure to check the absence of remaining PM vascularization.

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**Institutional Review Board Statement:** The study was conducted according to the guidelines of the Declaration of Helsinki, and approved by the Institutional Review Board of Institut Paoli-Calmettes (the IRB number of this study is: PRAM-I-IPC 2018-013, approval date: 4 July 2018).

**Informed Consent Statement:** Informed consent was obtained from all subjects involved in the study.

**Data Availability Statement:** The data presented in this study are available on request from the corresponding author. The data was generated from clinical history of patients therefore they are not publicly available.

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**Conflicts of Interest:** The 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.

**References**

1. Ballarin, R.; Spaggiari, M.; Cautero, N.; Ruvo, N.D.; Montalti, R.; Longo, C.; Pecchi, A.; Giacobazzi, P.; Marco, G.D.; D’Amico, G.; et al. Pancreatic Metastases from Renal Cell Carcinoma: The State of the Art. *World J. Gastroenterol.* 2011, 17, 4747. [CrossRef]

2. Gravis, G.; Chanez, B.; Derosa, L.; Beuselinck, B.; Barthelemy, P.; Laguerre, B.; Branchet, P.-E.; Joly, F.; Escudier, B.; Harrison, D.J.; et al. Effect of Glandular Metastases on Overall Survival of Patients with Metastatic Clear Cell Renal Cell Carcinoma in the Antiangiogenic Therapy Era. *Urol. Oncol.* 2016, 34, 167.e17–167.e23. [CrossRef] [PubMed]

3. Beuselinck, B.; Oudard, S.; Rixe, O.; Wolter, P.; Blessius, A.; Ayllon, J.; Elaidi, R.; Schöffski, P.; Barrascout, E.; Morel, A.; et al. Negative Impact of Bone Metastasis on Outcome in Clear-Cell Renal Cell Carcinoma Treated with Sunitinib. *Ann. Oncol.* 2011, 22, 794–800. [CrossRef]

4. Flippot, R.; Dalban, C.; Laguerre, B.; Borchiellini, D.; Gravis, G.; Négrier, S.; Chevreau, C.; Joly, F.; Geoffrois, L.; Ladoire, S.; et al. Safety and Efficacy of Nivolumab in Brain Metastases From Renal Cell Carcinoma: Results of the GETUG-AFU 26 NIVOREN Multicenter Phase II Study. *J. Clin. Oncol.* 2019, 37, 2008–2016. [CrossRef] [PubMed]

5. McKay, R.R.; Kroeger, N.; Xie, W.; Lee, J.-L.; Knox, J.J.; Bjarnason, G.A.; MacKenzie, M.J.; Wood, L.; Srinivas, S.; Vaishampayan, U.N.; et al. Impact of Bone and Liver Metastases on Patients with Renal Cell Carcinoma Treated with Targeted Therapy. *Eur. Urol.* 2014, 65, 577–584. [CrossRef]

6. Turajlic, S.; Xu, H.; Litchfield, K.; Rowan, A.; Chambers, T.; Lopez, J.J.; Nicol, D.; O’Brien, T.; Larkin, J.; Horswell, S.; et al. Tracking Cancer Evolution Reveals Constrained Routes to Metastases: TRACERx Renal. *Cell 2018*, 173, 581–594.e12. [CrossRef]

7. Singla, N.; Xie, Z.; Zhang, Z.; Gao, M.; Youssuf, Q.; Onabolu, O.; McKenzie, T.; Tcheyuyap, V.T.; Ma, Y.; Choi, J.; et al. Pancreatic Tropism of Metastatic Renal Cell Carcinoma. *JCI Insight* 2020, 5, e134564. [CrossRef]

8. Dabezati, S.; Marconi, L.; Hofmann, F.; Stewart, F.; Lam, T.B.L.; Canfield, S.E.; Staehler, M.; Powles, T.; Ljungberg, B.; Bex, A. Local Treatments for Metastases of Renal Cell Carcinoma: A Systematic Review. *Lancet Oncol.* 2014, 15, e549–e561. [CrossRef] [PubMed]

9. Tosioian, J.J.; Cameron, J.L.; Allaf, M.E.; Hruban, R.H.; Nahime, C.B.; Pawlik, T.M.; Pierrozzi, P.M.; Reddy, S.; Wolfgang, C.L. Resection of Isolated Renal Cell Carcinoma Metastases of the Pancreas: Outcomes from the Johns Hopkins Hospital. *J. Gastrointest. Surg.* 2014, 18, 542–548. [CrossRef]

10. Santoni, M.; Conti, A.; Partelli, S.; Porta, C.; Sternberg, C.N.; Procopio, G.; Bracarda, S.; Basso, U.; Giorgi, U.D.; Derosa, L.; et al. Surgical Resection Does Not Improve Survival in Patients with Renal Metastases to the Pancreas in the Era of Tyrosine Kinase Inhibitors. *Ann. Surg. Oncol.* 2014, 22, 2094–2100. [CrossRef]

11. Zerbi, A.; Ortolano, Ê.; Balzano, G.; Borri, A.; Beneduce, A.A.; Carlo, V.D. Pancreatic Metastasis From Renal Cell Carcinoma: Which Patients Benefit from Surgical Resection? *Ann. Surg. Oncol.* 2008, 15, 1161–1168. [CrossRef]

12. Yuasa, T.; Inoshita, N.; Saito, A.; Yamamoto, S.; Urakami, S.; Masuda, H.; Fujii, Y.; Fukui, I.; Ishikawa, Y.; Yonese, J. Clinical Outcome of Patients with Pancreatic Metastases from Renal Cell Cancer. *BMC Cancer* 2015, 15, 46. [CrossRef]
13. Sperti, C.; Moletta, L.; Patanè, G. Metastatic Tumors to the Pancreas: The Role of Surgery. *World J. Gastrointest. Oncol.* 2014, 6, 381–392. [CrossRef]

14. Benhaim, R.; Oussoultzoglou, E.; Saeedi, Y.; Mouracade, P.; Bachellier, P.; Lang, H. Pancreatic Metastasis from Clear Cell Renal Cell Carcinoma: Outcome of an Aggressive Approach. *Urology* 2015, 85, 135–140. [CrossRef]

15. Alvarez-Sánchez, M.-V.; Napoléon, B. Review of Endoscopic Radiofrequency in Biliopancreatic Tumours with Emphasis on Clinical Benefits, Controversies and Safety. *World J. Gastroenterol.* 2016, 22, 8257–8270. [CrossRef] [PubMed]

16. Chu, K.F.; Dupuy, D.E. Thermal Ablation of Tumours: Biological Mechanisms and Advances in Therapy. *Nat. Rev. Cancer* 2014, 14, 199–208. [CrossRef] [PubMed]

17. Barthet, M.; Giovannini, M.; Lesavre, N.; Boustiere, C.; Napoleon, B.; Koch, S.; Gasmi, M.; Vanbiervliet, G.; Gonzalez, J.-M. Endoscopic Ultrasound-Guided Radiofrequency Ablation for Pancreatic Neuroendocrine Tumors and Pancreatic Cystic Neoplasms: A Prospective Multicenter Study. *Endoscopy* 2019, 51, 836–842. [CrossRef] [PubMed]

18. Elias, D.; Baton, O.; Sideris, L.; Lasser, P.; Pocard, M. Necrotizing Pancreatitis after Radiofrequency Destruction of Pancreatic Tumours. *Eur. J. Surg. Oncol.* 2004, 30, 85–87. [CrossRef]

19. Rochigneux, P.; Nault, J.-C.; Mallet, F.; Chretien, A.-S.; Barget, N.; Garcia, A.J.; Del Pozo, L.; Bourcier, V.; Blaise, L.; Grando-Lemaire, V.; et al. Dynamic of Systemic Immunity and Its Impact on Tumor Recurrence after Radiofrequency Ablation of Hepatocellular Carcinoma. *OncoImmunology* 2019, 8, 1615818. [CrossRef] [PubMed]

20. Zerbini, A.; Pilli, M.; Schiantchi, C.; Molinari, A.; Negri, E.; Cerioni, S.; Fagnoni, F.; Soliani, P.; Ferrari, C.; et al. Radiofrequency Thermal Ablation of Hepatocellular Carcinoma Liver Nodules Can Activate and Enhance Tumor-Specific T-Cell Responses. *Cancer Res.* 2006, 66, 1139–1146. [CrossRef] [PubMed]