Incidence and clinical presentation of adrenocortical carcinoma (ACC)

Incidental discovery of adrenal tumors is common and occurs in up to 7% of patients who undergo cross-sectional imaging for other reasons (1). The vast majority of these adrenal “incidentalomas” are benign, nonfunctioning adenomas. ACC itself is quite rare with an incidence of only 1–2 cases per million per year (2). ACC occurs slightly more often in women compared to men, with a ratio of approximately 1.5:1 (3). Though it has been diagnosed at any age, peak incidences occur in both the first half of the first decade of life, and the fourth and fifth decades of life (4,5). Autonomous steroid production, such as hypercortisolism (Cushing’s syndrome) or androgen excess, is present in around 60% of patients with ACC (4-7). The diagnosis of ACC is often presumptive based on imaging (CT or MRI) and hormonal workup and confirmed post-operatively on surgical pathology (3). Biopsy of lesions concerning for ACC is not recommended due to the potential for tumor spillage.

Imaging characteristics of ACCs

In assessing adrenal tumors, certain imaging characteristics are helpful in stratifying risk of malignancy. On CT imaging, ACCs are typically seen as large, heterogeneous suprarenal masses. Increasing tumor size is the most
commonly cited risk factor for ACC (8). Whereas only 2% of adrenal incidentalomas ≤4 cm or less are ACC, one quarter of incidental adrenal tumors larger than 6 cm have been found to be ACC (5,9). One large study found that the risk of an adrenal cortical tumor being ACC is 10% when ≥4 cm, 19% when ≥6 cm, and 47% when ≥8 cm (10). Another study found 31/37 (84%) of adrenal cortical lesions ≥8 cm to be ACC, while only 6/37 (16%) were benign adenomas (P<0.01) (11). Most surgeons agree that non-functioning tumors less than 4 cm can be observed while tumors greater than 6 cm (except for asymptomatic myelolipomas) should be resected regardless of hormonal production status. The management of tumors between 4–6 cm is less clear and shared-decision making should be employed with patients to determine the possible need for resection (12).

Adrenal mass density characteristics on CT scan can also be used to help differentiate adenomas from suspected ACCs. The vast majority of adrenal masses measuring less than 10 Hounsfield units on non-contrast imaging represent adenomas (sensitivity 71%; specificity of 98% for adenoma) (13). Though most ACCs measure greater than 10 Hounsfield units on non-contrast imaging, use of this criterion alone to diagnose ACC results in a high false positive rate, with reported specificities as low as 58% (5,14,15). ACCs can further be distinguished from benign adrenal adenomas by their slower contrast washout (14). Unlike adenomas, which demonstrate both rapid uptake and washout of contrast on CT imaging, ACCs and other non-adenomatous lesions have a delayed contrast washout. In general, thresholds of 60% contrast enhancement washout and 40% relative contrast enhancement washout have been able to distinguish adenomas from non-adenomas, including ACC, with 92–100% sensitivity and 95–100% specificity (14,16-18). It should be noted that most of these imaging analyses have compared washout characteristics between adenomas and non-adenomas, and not explicitly adrenal adenomas from ACCs. One study examining washout characteristics of ACCs only found that out of 17 patients with ACCs, 71% had ≤60% contrast enhancement washout and 82% had ≤40% relative contrast enhancement washout using a 15-min delay (14). Therefore, some caution should be used relying solely on washout characteristics to “rule out” ACC.

Calcification and necrosis are common features of ACCs. Calcifications are observed in up to 37% of patients with ACC (14). They are typically located centrally and can be identified on routine pre-operative imaging (5,14). Moreover, ACCs often have central areas of hemorrhage and necrosis, and therefore demonstrate more peripheral than central enhancement with contrast administration (19).

Since ACCs are commonly large at the time of diagnosis, invasion of adjacent organs and the venous system is not unusual. In an imaging-focused retrospective study, seven out of 41 patients with ACC (17%) were found to have invasion into adjacent organs: two with liver invasion, two with kidney invasion, two with combined liver-kidney invasion, and one with combined pancreas, stomach, and diaphragm invasion (14). In the same retrospective study, 6 out of 41 patients (15%) were found to have ACC extension and associated thrombus in the inferior vena cava (IVC) (14).

### Staging in ACC

Staging for ACC is based on the European Network for the Study of Adrenal Tumors (ENSAT) and American Joint Committee on Cancer (AJCC) staging system (Table 1) (20,21). ACC is classified into one of four stages. Broadly, stages I and II disease are confined to the adrenal gland, with a cutoff of 5 cm differentiating Stage I from Stage II disease. Stage III and IV ACC represent lymph node

| Table 1 ENSAT/AJCC Staging System and Associated 5-Year Survival Data | 5-Year Survival |
| --- | --- |
| Stage I | T1 M0 N0 82% |
| Stage II | T2 M0 N0 61% |
| Stage III | T1–2 or T3–4 M0 N0–1 50% |
| Stage IV | T any M0 N any M1 13% |

1, Staging classification and 5-year survival data from Fassnacht et al. (20). ‘Survival’ refers to disease-specific survival. T1, tumor ≤5 cm; T2, tumor >5 cm; T3, tumor invasion of surrounding tissue; T4, tumor invasion of adjacent organs or tumor thrombus in vena cava or renal vein; N0, no lymph node positivity; N1, lymph node positivity; M0, no distant metastases; M1, distant metastases.
positivity (stage III or IV), local invasion (stage III or IV), or metastatic disease (stage IV). This staging system has been validated using data derived from 573 patients in 16 Surveillance, Epidemiology and End Results (SEER) registries (22). This validation confirmed the prognostic superiority of ENSAT over its predecessor, the International Union Against Cancer (UICC) staging system, which failed to differentiate prognostic implications of stage II versus stage III ACC (20,22).

**Pre-operative workup for suspected ACC**

The European Society of Endocrine Surgeons (ESES) and European Network for the Study of Adrenal Tumours (ENSAT) have published guidelines for the pre-operative management of adrenal masses that are suspected to be ACC (19). Although the low overall quality of evidence due to the lack of prospective or randomized studies is acknowledged, the authors compiled consensus recommendations in light of the available evidence.

In taking a history, patients should be evaluated for symptoms associated with excess hormone production (5,7,23) and symptomatic compression due to the large size of a mass (5,7,23,24). A past medical history of malignancy is important since some adrenal tumors may represent metastases from another primary cancer. Lastly, a thorough family history should be obtained to assess for familial cancers and help guide the need for genetic assessment (25-28).

A full electrolyte and hormonal evaluation are recommended. Plasma and urinary metanephrines are checked to assess for the possibility of a pheochromocytoma, which carries its own unique clinical and operative management considerations (19). Additionally, the guidelines recommend biochemical assessment of steroid production (sex steroids and their precursors, aldosterone, and cortisol) as functional status of the tumor may necessitate post-operative steroid replacement. A prospective study is underway to determine the utility of 24-hour urine steroid profiling using mass spectrometry (29).

Pre-operative imaging recommendations for adrenal tumors are intended to identify those that are more likely to be malignancies, as well as to guide operative management. For suspected ACC, contrast-enhanced CT of the chest and abdomen within 6 weeks of surgery is recommended to identify characteristics of the tumor, lymph node involvement, and to assess for intravascular thrombus or invasion of the tumor into local structures. The guidelines also include gadolinium contrast-enhanced MRI as an essential component of an adrenal tumor workup if its risk of malignancy cannot be determined on CT alone. MRI is also more sensitive than CT for detection of tumor thrombus, vascular invasion, or liver involvement (5,30).

Advanced imaging studies to accurately stage the disease and evaluate for distant metastasis can be used. $[^{18}\text{F}]$ fluoro-deoxyglucose- (FDG-) PET is recommended within 6 weeks of adrenalectomy for adrenal tumors suspected to be ACC. Adrenal tumors that demonstrate FDG-avidity include ACC (31,32), pheochromocytomas (33), and some adrenal adenomas (34), though adenomas may have lower uptake when uptake is present. The specificity of FDG-PET for ACC in a series of 77 patients was 88% when using a maximum standardized uptake value cutoff of 1.34 (31); therefore, it is best utilized to stage and follow the trajectory of the disease after surgical intervention. $[^{11}\text{C}]$ metomidate and $[^{123}\text{I}]$iodometomidate are more specific for adrenocortical cells but they are limited by clinical availability (35) and an inability to distinguish between benign and malignant lesions.

**Survival outcomes in patients with ACC**

The rarity of this malignancy complicates precise assessment of outcomes. Prognosis is correlated with surgical margin status following resection and stage of disease at the time of diagnosis (6,36-44). In one study, recurrence-free survival (RFS) was as low as 18% at median follow-up of 88 months, with a median time to recurrence of 13 months (38). Surgical margin status is a critical factor associated with recurrence since R0 resection of ACC confers a decreased risk of recurrent disease compared to R1 or R2 resection (39,45-50). A review of 3,982 patients with ACC from the National Cancer Data Base found adjusted hazard ratios for 5-year survival of 1.81 (95% CI, 1.44–2.27) for positive microscopic margins and 2.06 (95% CI, 1.74–2.43) for positive macroscopic margins when compared to R0 resection status (39). Five-year overall survival (OS) rates range from 15% to 60%, with as high as 84% 5-year OS for stage I disease compared to only 18% survival for stage IV disease (20,39,46,51-56). Interestingly, the US ACC study group recently reported that patients with incidentally discovered ACC may present at an earlier stage compared to patients with functional and symptomatic ACC leading to potentially improved recurrence free survival for incidentally discovered tumors (57). Nonetheless, ACC is often discovered at advanced stages with approximately 70% of tumors stage II or greater at the time of diagnosis.
Advanced stage at diagnosis and a lack of effective systemic therapy options contribute to its poor overall prognosis (36,58,59).

**Surgical management of ACC**

Open adrenalectomy (OA) is the gold standard treatment for suspected ACC when technically feasible (19,58). The past 20 years have witnessed the advent and rapid expansion of minimally invasive surgery in surgical oncology. In order of importance, the priorities during surgery for suspected ACC are (I) oncologic control and (II) application of a minimally invasive approach. The first report of laparoscopic adrenalectomy (LA) for ACC was published in 1993 (60). Other early case reports and small case series began reporting on the role of laparoscopy in adrenalectomy for ACC in the late 1990s. These small studies and single patient reports yielded results that were split between optimism for a new, minimally invasive approach to a historically open procedure (61-65), and apprehension due to concern for increased risks of locoregional recurrence and peritoneal carcinomatosis (66-72).

**Studies in which the data favor open adrenalectomy**

Despite initial enthusiasm for LA, a number of studies have published data favoring an OA over LA in patients with suspected ACC (Table 2) (70,73-80). The first of these analyzed results from 159 patients with ACC, though only 6 underwent LA against 153 undergoing OA (70). Follow-up data were available for 133 patients in the OA group, 86% of whom had experienced recurrence at a median follow-up of 28 months. At follow-up, all six patients who had undergone LA had experienced recurrence. Additionally, whereas only 11 of 133 patients (8%) in the OA group had peritoneal carcinomatosis present at the time of initial failure, it was present in five out of six patients in the LA group (83%), leading the authors to conclude that OA is superior to LA for suspected ACC (70).

Data from five series were published between 2010 and 2013. In 2010, Lebougleux et al. published retrospective data on 64 patients, 58 of whom underwent OA versus 6 who underwent LA (75). While the sample of patients undergoing LA was low, LA once again portended an increased risk of peritoneal carcinomatosis (67% in the LA group compared to a 27% in the OA group at four years after adrenalectomy). After analysis of other metrics such as tumor size and stage, LA was found to be the only variable associated with peritoneal carcinomatosis, with a 4-year hazard ratio (HR) of 3.8 (95% CI, 1.2-12.3).

Another study published in 2010, from Miller et al., compared retrospective outcomes of 88 patients with stage I-III ACC (76). Slightly more LA cases were included in this series, with 71 patients having undergone OA versus 17 undergoing LA. This study found that although tumor size was unbalanced between groups and were more favorable in the LA group (median tumor size was 7.0 cm in LA group versus 12.3 cm in OA group), outcomes favored OA nonetheless. Significant findings included a longer mean time to local recurrence (19.2±7.5 versus 9.6±14 months, P=0.005) and a lower positive margin or intraoperative tumor spill rate (18% versus 50%, P=0.01) in the OA group compared to the LA group.

In 2012, Miller et al. published an update on these 88 patients and included 68 new patients, with stages I-III ACC who had undergone either LA or OA (77). Several important outcomes in this study favored an open over laparoscopic approach to adrenalectomy for suspected ACC. Positive margins, an important prognostic indicator for patients after adrenalectomy, were more likely to be present after LA compared to after OA (30% versus 16%, P=0.04). Recurrence in the tumor bed occurred faster in patients undergoing LA compared to patients undergoing OA (mean time: 11.7 versus 30.5 months, P=0.002). This pattern persisted for recurrence in any distant location (mean time: 17.6 after LA vs. 30.5 months after OA, P=0.002). Mean OS was significantly longer in patients with stage II ACC who underwent OA compared to LA (103.1 versus 50.9 months, P=0.002). A critical finding from Miller et al. was the discovery that 31% of clinical stage II patients were upstaged to stage III based on post-operative pathology. This finding has implications for those advocating tumor stage I or II as a criterion for choosing LA over OA for ACC (24,81-85), as evidence suggest that up to one in three patients selected for LA based on their clinical stage II are understaged, and may be inappropriate candidates for LA.

In 2013, Mir et al. reported retrospective data on 44 patients, 26 of whom had undergone OA, and 18 of whom had undergone LA for stages I-IV ACC (78). Similar to the 2010 Miller study, median tumor size was larger in the OA group compared to the LA group (13 versus 7 cm, P=0.001) (76,78). In addition, patients in the OA group had more advanced tumors at baseline when compared to the LA group (62% stages III-IV versus 11% stages III-IV, P=0.001). Despite the poorer pre-operative tumor profiles...
Table 2 Retrospective analyses in favor of open adrenalectomy for patients with adrenocortical carcinoma

| Author       | Year | Study details† | Important results                                                                 | Author conclusions/recommendations                                                                                                                                                                                                 |
|--------------|------|----------------|------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Gonzalez (70) | 2005 | N=159          | Recurrence rate (median follow-up 28 months) 115/133 (86%) after OA vs. 6/6 (100%) after LA | Peritoneal carcinomatosis is common after LA for ACC, and OA is superior to LA for adrenal masses that may be ACC                                                                                                                |
|              |      | 153 OA, 6 LA   | PC at initial failure: 11/133 (8%) after OA vs. 5/6 (83%) after LA                  |                                                                                          |
|              |      | Surgeries performed at multiple locations |                                                                                           |                                                                                          |
| Cooper (73)  | 2013 | N=302          | Median tumor size: Smaller in LA vs. OA groups (8.0 cm in LA vs. 12.0 cm in OA at outside hospital vs. 12.3 cm for OA at index hospital) | Patients with suspected ACC should undergo OA as opposed to LA when controlling for clinical stage (exact values not given)                                                                                       |
|              |      | 256 OA (210 at outside centers, 46 at index hospital), 46 LA | R0 margin status: 54.3% after LA vs. 44.3% after LA at outside hospital vs. 89.1% after OA at index hospital |                                                                                                                                                                                                                           |
|              |      | Surgeries performed at multiple centers | PC: 54.3% after LA vs. 27.6% after LA at outside hospital vs. 19.6% after OA at index hospital (P=0.006) |                                                                                                                                                                                                                           |
|              |      | Stages I–IV    | RFS and OS for OA were statistically significantly superior to LA                   |                                                                                                                                                                                                                           |
| Huynh (74)   | 2016 | N=423          | Patients in OA group were younger, had more advanced disease, and had larger tumors than LA group | As OS may decrease in some patients with ACC treated with LA, “caution should still be used in selecting LA for surgical treatment of ACC.”                                                                                      |
|              |      | OA 286, LA 137 | Positive margin rate: statistically significant increase in LA (54.6%) vs. OA (21.7%) in T3 tumors only (P=0.0009) |                                                                                                                                                                                                                           |
|              |      | Data obtained from National Cancer Center Database | OA vs. LA was not associated with OS for patients in overall cohort; positive margin was the only factor associated with decreased OS |                                                                                                                                                                                                                           |
|              |      | Stages I–III   | OS in patients with T2 disease: lower in LA vs. OA (P=0.04)                         |                                                                                                                                                                                                                           |
|              |      | On MVA, LA was independent risk factor for death (HR 1.86, 95% CI, 1.02 to 3.38; P=0.04) |                                                                                                                                                                                                                           |
| Leboulleux (75) | 2010 | N=64           | LA (67%, 95% CI, 30–90%) was associated with greater 4-year risk of development of PC vs. OA (27%, 95% CI, 15–44%); P=0.016 | There is an increased risk of PC after LA for ACC, though it is unclear if this is due to the surgical approach or surgical inexperience with ACC                                                                                       |
|              |      | 58 OA, 6 LA    | No other factors (e.g., tumor size, tumor stage) were associated with PC            |                                                                                                                                                                                                                           |
|              |      | Single center  |                                                                                   |                                                                                                                                                                                                                           |
|              |      | Stages I–IV    |                                                                                   |                                                                                                                                                                                                                           |
Table 2 (continued)

| Author      | Year | Study details | Important results                                                                 | Author conclusions/recommendations |
|-------------|------|---------------|-----------------------------------------------------------------------------------|-------------------------------------|
| Miller (76) | 2010 | N=88          | Median tumor size: smaller in LA (7.0 cm) vs. OA (12.3 cm)                        | LA should not be attempted for tumors that may be ACC |
|             |      |               | Disease recurrence: 63% after LA vs. 65% after OA                                 |                                     |
|             |      |               | Surgeries performed at multiple locations                                         |                                     |
|             |      |               | Mean time to first recurrence: 9.6±14 months after LA vs. 19.2±37.5 months after OA (P<0.005) |                                     |
|             |      |               | Stages I–III                                                                      |                                     |
|             |      | 71 OA, 17 LA  | Positive margin or intraoperative tumor spill rate: 50% in LA group vs. 18% in OA group (P=0.01) |                                     |
|             |      |               | Local recurrence: 25% after LA vs. 20% after OA (P=0.23)                          |                                     |
| Miller (77) | 2012 | N=156         | Positive margins: 30% after LA vs. 16% after OA (P=0.04)                          | Patients with ACC should undergo OA, and “intraoperative evaluation is insensitive for the detection of stage III tumors” |
|             |      |               | 110 OA, 46 LA                                                                    |                                     |
|             |      |               | Mean OS in stage II ACC: 50.9 months after LA vs. 103.1 months after OA (P=0.002) |                                     |
|             |      | Surgeries performed at multiple locations                                         |                                     |
|             |      |               | Mean time to tumor bed or peritoneal cavity recurrence: 11.7 months after LA vs. 30.5 months after OA (P=0.002) |                                     |
|             |      | Stages I–III | Mean time to distant recurrence: 17.6 months after LA vs. 52.9 months after OA (P=0.001) |                                     |
|             |      |               | 35/113 (31%) of all presumed stage II patients were upstaged to stage III post-operatively |                                     |
|             |      |               | Local recurrence: 25% after LA vs. 20% after OA (P=0.23)                          |                                     |
|             |      |               | Mean time to distant recurrence: 17.6 months after LA vs. 52.9 months after OA (P=0.001) |                                     |
| Mir (78)    | 2013 | N=44          | OA group had larger, more advanced tumors                                         | It is difficult to assess differences in outcomes given the rarity of ACC, but this data “suggests superiority for OA in the management of ACC” |
|             |      |               | 26 OA, 18 LA                                                                      |                                     |
|             |      | Single center  | 2-year OS: 54% after OA vs. 58% after LA (P=0.6)                                 |                                     |
|             |      |               | Non-statistically significant benefit was found for OS in OA vs. LA after adjusting for clinical stage |                                     |
| Wu (79)     | 2018 | N=44          | Mean LOS: shorter after LA (6±2 days) vs. OA (9±4 days); (P=0.002)                | OA should remain the standard surgical approach. Compared to OA, LA may not yield comparable outcomes for patients with localized ACC. The authors extend this recommendation even to patients with tumor size <10 cm and no evidence of extra-adrenal invasion |
|             |      |               |                                                                                   |                                     |

Table 2 (continued)
Table 2 (continued)

| Author      | Year | Study details | Important results                                                                                                                                                                                                 | Author conclusions/recommendations                                                                 |
|-------------|------|---------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------|
| OA 23, LA 21 |      |               | Overall disease recurrence: no difference after LA (52%) vs. after OA (52%); P=0.989                                                                                                                              |                                                                                                                                                                |
| Single center |     |               | Local recurrence: 42% after LA vs. 22% after OA (P=0.13)                                                                                                                                                           |                                                                                                                                                                |
| Stages I–II, tumor size ≤10 cm |      |               | Distant metastases: 24% after LA vs. 30% after OA (P=0.622)                                                                                                                                                         |                                                                                                                                                                |
|             |      |               | PC rate: 24% after LA vs. 4.3% after OA (P=0.15)                                                                                                                                                                 |                                                                                                                                                                |
|             |      |               | Mean time to recurrence: no difference between LA (25±22 months) vs. OA (22±24 months); P=0.564                                                                                                               |                                                                                                                                                                |
|             |      |               | Mean time to local recurrence and PC: faster after LA (40±8 months) vs. OA (79±9 months), P=0.048                                                                                                               |                                                                                                                                                                |
|             |      |               | 5-year OS: 43% after OA vs. 47% after LA (P=0.635)                                                                                                                                                              |                                                                                                                                                                |
|             |      |               | 5-year RFS: 36% after OA vs. 39% after LA (P=0.802)                                                                                                                                                              |                                                                                                                                                                |
| Zheng (80)  | 2018 | N=42          | Mean tumor size: larger in OA group (10.1±3.6 cm) vs. LA group (6.3±2.2 cm), P=0.001                                                                                                                             | Even though short-term outcomes like LOS and EBL favor LA, OA is still superior to LA for ACC given DFS benefit and risk of local recurrence after LA |
| OA 22, LA 20 |      |               | Median operative time: longer in OA group (175 min) vs. LA group (130 min); P=0.004                                                                                                                              |                                                                                                                                                                |
| Single center |     |               | Median EBL: Greater in OA group (800 mL) vs. LA (70 mL), P=0.001                                                                                                                                             |                                                                                                                                                                |
| Stages I–III |      |               | Median LOS: Longer in OA group (9.5 d) vs. LA group (7.0 d), P=0.018                                                                                                                                         |                                                                                                                                                                |
|             |      |               | Mean DFS: Longer in OA group (44.8±35.1 months) vs. LA group (17.5±10.4 months), P=0.023                                                                                                                       |                                                                                                                                                                |
|             |      |               | 2-year DFS: higher in OA group (61.1%) vs. LA group (21.4%), P=0.036                                                                                                                                     |                                                                                                                                                                |
|             |      |               | Difference in DFS at 1 and 3 years not statistically significant between OA and LA groups                                                                                                                         |                                                                                                                                                                |
|             |      |               | Locally recurrent disease at first recurrence: 62% in OA group vs. 100% in LA group (P=0.03)                                                                                                                           |                                                                                                                                                                |

†, in this column, “N” refers to the total number of patients in each study undergoing either laparoscopic or open adrenalectomy, not the total number of patients included in the study. OA, open adrenalectomy; LA, laparoscopic adrenalectomy; PC, peritoneal carcinomatosis; ACC, adrenocortical carcinoma; CI, confidence interval; OS, overall survival; RFS, recurrence-free survival; DFS, disease-free survival; MVA, multivariate analysis; LOS, length of stay; EBL, estimated blood loss.
in the OA group, outcomes between the two groups were similar, with no significant differences in overall 2-year overall or RFS. When adjustments were made for clinical stage, overall and disease-free survival benefits trended in favor of OA, but did not reach statistical significance (HR for recurrence 0.4, 95% CI, 0.2–1.2, P=0.099; HR for OS 0.5, 95% CI, 0.2–1.2, P=0.122).

A larger retrospective analysis published by Cooper et al. in 2013 found long-term outcomes that favored OA over LA for patients with ACC (73). This study evaluated 302 patients with all stages of ACC. Two hundred ten underwent OA at outside centers, 46 underwent OA at the index hospital, and 46 underwent LA (all at outside centers). Median tumor size was larger in the OA versus LA groups (8.0 cm in LA group, 12.0 cm in OA group from outside hospitals, and 12.3 cm in OA group from index hospital). In this study, peritoneal carcinomatosis occurred in 54.3% of patients after undergoing LA, 27.6% after OA at an outside hospital, and 19.6% after OA at the index hospital (P=0.006). Additionally, RFS and OS were superior to LA when the authors controlled for clinical stage (P<0.0001 for both). Similar to Miller et al. (77), the authors of this paper concluded that patients should undergo OA if they have an adrenal tumor suspected to be ACC (73).

The largest study comparing open versus laparoscopic adrenalectomy for suspected ACC was published by Huynh et al. in 2016 (74). Records of 423 patients who had undergone OA (286 patients) or LA (137 patients) for stages I-III ACC were identified from the National Cancer Center Database. Despite patients in the OA group having larger, more advanced tumors compared to the LA group, LA was identified as an independent risk factor for death on multivariate analysis (HR 1.86, 95% CI, 1.02–3.38; P=0.04). For patients with T3 tumors, the positive margin rate was significantly higher in the LA group compared to the OA group (54.6% versus 21.7%, P=0.0009). Although LA was associated with a significant increase in positive margin rate, surgical approach was not directly associated with OS in the overall cohort. For patients with T2 tumors, OS was better for OA compared to LA (P=0.04). The authors concluded that because OS may decrease in some patients with ACC treated with LA, “caution should still be used in selecting LA for surgical treatment of ACC” (74).

Two more retrospective studies were published in 2018. Zheng et al. reported retrospective data on 42 patients (22 OA versus 20 LA) with stages I–III ACC (80). Similar to previous studies, median tumor size was significantly larger in the OA group compared to the LA group. In this study, short-term metrics were evaluated in addition to longer-term outcomes. Although the short-term data, such as intra-operative blood loss, operative time, and length of stay all favored LA, long-term data favored OA. Mean DFS was significantly longer in the OA group compared to the LA group (44.8±35.1 versus 17.5±10.4 months, P=0.023).

Later in 2018, Wu et al. published data on 44 patients who had undergone OA (23 patients) or LA (21 patients) for stages I-II ACC with tumor size less than 10 cm (79). Mean duration of post-operative hospital stay was shorter after LA versus OA by an average of 3 days (6 versus 9 days, P=0.002). Rates of local recurrence (42% versus 22%, P=0.13) and peritoneal carcinomatosis (24% versus 43%, P=0.15) trended in favor of OA but did not reach statistical significance. However, mean time to local recurrence and peritoneal carcinomatosis was significantly shorter in the LA group compared to the OA group (40±8 versus 79±9 months, P=0.048). Five-year overall (43% after OA versus 47% after LA, P=0.635) and risk-free survival (36% after OA versus 39% after LA, P=0.802) was similar between groups.

A meta-analysis published by Autorino et al. in 2016 examined nine retrospective case-controlled studies published between 2010 and 2014 (240 total patients underwent LA, 557 patients underwent OA) (86). No differences were found in metrics such as overall recurrence (relative risk 1.09, 95% CI, 0.83–1.43, P=0.52), time to recurrence (weight mean difference –8.2 months, 95% CI, –18.2–1.7 months, P=0.11), or cancer-specific mortality (odds ratio 0.68, 95% CI, 0.44–1.05, P=0.08) when comparing LA to OA. Peritoneal carcinomatosis occurred more often in the LA group compared to the OA group (RR 2.39; 95% CI, 1.41–4.04; P=0.001). The results of this meta-analysis led the authors to conclude that while LA has short-term benefits of shorter length of stay and recovery time, OA should be considered the standard of care in adrenal tumors suspected of being ACC, with LA reserved for carefully selected patients in centers with high expertise.

Studies in which the data favor laparoscopic adrenalectomy

In order for LA to supplant OA as standard of care for surgical resection of suspected ACC, it must demonstrate long-term oncologic outcomes comparable to OA while providing short-term benefits inherent to the minimally invasive approach. Case reports and case series from 1999 to 2005 detailed technical efficacy using laparoscopy to
treat ACC (27-31). Since then, results of some retrospective analyses directly comparing laparoscopic versus the open adrenalectomy have concluded that LA is a feasible option for patients with ACC, particularly in carefully selected cases (Table 3) (24,81-85,87,88).

In 2008, Kirschstein et al. published data on 12 patients who had undergone adrenalectomy for stages I–III ACC at a single institution (82). Seven underwent OA and 5 underwent LA. No recurrences were reported at the time of publication. The length of follow-up was not reported. Given the short-term benefits in the laparoscopic group (lower estimated blood loss and shorter length of stay), the authors deemed LA as a viable option for carefully selected patients with ACC based on short-term data.

Porpiglia et al. reported on 43 patients (25 OA, 18 LA) who underwent adrenalectomy for stages I–II ACC at multiple surgical centers (88). In this series, baseline demographics were similar between groups, with no statistically significant differences in median age (41.3 versus 47, P=0.16), tumor stage (88% stage I versus 80% stage II, P=0.4), or median tumor size (10.5 versus 9 cm, P=0.39) in the open versus laparoscopic groups. Recurrence occurred in 50% of patients undergoing LA compared to 64% of patients undergoing OA. Median RFS was similar in the OA group versus the LA group (18 versus 23 months, P=0.8). Mortality during follow-up trended in favor of LA (5%) compared to OA (28%). The authors concluded that LA was feasible in the management of ACC, but did not definitively confirm its efficacy.

Brix et al. analyzed data from a larger group of patients derived from a German registry of ACC patients undergoing OA or LA for stages I–III ACC less than 10 cm (81). Out of the 152 patients included in the study, 117 underwent OA and 35 underwent LA. The occurrence of peritoneal carcinomatosis was 3% in both groups. No differences between groups were seen regarding disease-specific survival (HR for death 0.98, 95% CI, 0.51–1.92, P=0.92 on MVA) and adjusted RFS (HR 0.91, 95% CI, 0.56–1.46, P=0.69). Importantly, 12 out of 35 patients undergoing LA required conversion to an open procedure, although the authors reported no differences in the outcomes of the converted patients compared to patients whose surgeries were entirely laparoscopic.

Lombardi et al. reported data from multiple centers in 2012 on 156 patients who underwent OA (n=126) or LA (n=30) for stages I–II ACC (84). Rates of local recurrence were similar after OA and LA (19% versus 21%), with rates of distant metastases trending in favor of LA (31% after OA, 17% after LA; P>0.05) over a mean follow-up time of 42±35 months. Mean time to recurrence was also similar between groups, and 5-year disease-free survival trended in favor of the LA group (38.3% after OA versus 58.2% after LA; P>0.05).

In 2013, Fossa et al. reported outcomes on 32 patients from a single institution who underwent OA (n=15) or LA (n=17) for stages I–III ACC. Short-term benefits favored the LA group including median operative time (150 for LA versus 230 min for OA, P=0.005), median number of perioperative transfusions (0 units for LA versus 3 units for OA, P=0.04), median length of stay (6 days after LA versus 13 days after OA, P=0.001), and median estimated blood loss (<400 during LA versus 1,700 mL during OA, P<0.001). Of note, median tumor size was significantly larger in the OA group (13.0 cm in the OA group versus 8.0 cm in the LA group, P=0.002), which could account for some of the short-term differences. R0 resection rates were similar in both groups (80% after OA versus 71% after LA). Recurrence rates were high and similar in both groups (88% after OA versus 80% after LA). Median progression-free survival approached statistical significance in favor of LA (15.2 months after LA versus 8.1 months after OA, P=0.057), although this was not derived from a multivariable model. The authors proposed that because rates of early recurrence were high and similar regardless of the surgical approach, short-term benefits of LA such as length of stay and intraoperative blood loss should factor into decision-making.

Donatini and colleagues evaluated outcomes from patients who had undergone LA (n=13) or OA (n=21) for stages I–II ACC less than 10 cm at a single center (24). They reported R0 resection status in all patients, and found no differences in overall survival (81% versus 85%, P=0.634), recurrence rate (24% versus 31%, P=0.655), or disease-free survival (47 versus 46 months, P=0.893) between OA and LA groups at comparable median follow-up times. Reported short-term outcomes included a significantly shorter length of stay after LA, and similar surgical complication rates between the two groups (8% in LA group versus 14% in OA group, P=0.387).

Maurice et al. used the National Cancer Database to compare outcomes for patients with stages I–IV ACC undergoing OA (n=320) versus minimally invasive adrenalectomy (MIA; 131 laparoscopic, 30 robotic) (85). Benefits of a minimally invasive approach included shorter median length of stay (LOS) (3 versus 6 days, P<0.01) and a lower re-admission rate (4.4% versus 8.8%, P=0.08).
| Author       | Year | Study Details† | Important Results                                                                 | Author Conclusions/ Recommendations                                                                 |
|-------------|------|----------------|----------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------|
| Brix (81)   | 2010 | N=152          | DSS: No difference between LA and OA (HR for death 0.98; 95% CI, 0.51–1.92; P=0.92 on MVA) | "LA by an experienced surgeon is not inferior to OA" if tumor is ≤10 cm and ACC is localized       |
|             |      |                | 117 OA, 35 LA                                                                     | RFS: No difference between LA and OA (HR 0.91; 95% CI, 0.56–1.46; P=0.69)                       |
|             |      |                | All tumors ≤10 cm                                                                 | 12/35 patients in LA group required open conversion                                             |
|             |      |                | Data derived from German ACC Registry                                             | PC: 3% in both groups                                                                             |
|             |      |                | Stages I–III                                                                      |                                                                                                |
| Kirshtein (82) | 2008 | N=12           | No recurrence reported in either group                                             | LA is “feasible and safe” in carefully selected patients based on short-term data; long-term follow-up is still lacking |
|             |      |                | 7 OA, 5 LA                                                                        | Favorable mean EBL and LOS in LA (200 mL, 2 days) vs. OA (500 mL, 7 days)                        |
|             |      |                | Single center                                                                      | Larger mean tumor size in OA vs. LA groups (4 vs. 8 cm)                                         |
|             |      |                | Stages I–III                                                                      |                                                                                                |
| Lee (83)    | 2017 | N=201          | 30-day morbidity: no difference between MIA (29.3%) and OA (30.9%) groups (P=0.839) | MIS may be considered for ACC when tumor size is ≤10 cm; OA should still be performed for tumors ≥10 cm, or when there is local invasion or lymph node involvement |
|             |      |                | OA 154, MIA 47                                                                     | Rates of MIA were similar across the 13 sites included in the study                               |
|             |      |                | Multi-center                                                                      | Intraoperative tumor rupture: no difference between MIA (12.2%) and OA (9.4%), P=0.612         |
|             |      |                | ENSAT staging not used in study                                                    | R0 status: no difference between MIA (77.0%) vs. OA (72%), P=0.50                               |
|             |      |                |                                                                                  | On MVA, EBL (HR 1.013, 95% CI, 1.001–1.026, P=0.038) and T-stage (HR 2.102, 95% CI, 1.106–4.348, P=0.045) were the only predictive factors of OS; surgical approach was not (HR for OA vs. MIA 2.152, 95% CI, 0.601–7.714, P=0.239) |
| Lombardi (84) | 2012 | N=156          | Local recurrence rate: 19% after OA, 21% after LA                                  | With regard to oncologic outcomes, LA is not inferior to OA in stage I–II ACC; the authors note that OA is still “mandatory” in stage III/IV ACC |
|             |      |                | 126 OA, 30 LA                                                                     | Distant metastases rate: 31% after OA, 17% after LA (P= NS)                                      |
|             |      |                | Multi-center                                                                      | Mean time to recurrence: 27±27 months after OA, 29±33 months after LA (P=NS)                    |
|             |      |                | Stages I–II                                                                        | 5-year DFS: 38.3% after OA vs. 58.2% after LA (P= NS)                                          |
|             |      |                |                                                                                  | 5-year OS: 48% after OA vs. 67% after LA (P= NS)                                               |

Table 3 (continued)
| Author | Year | Study Details† | Important Results | Author Conclusions/ Recommendations |
|--------|------|----------------|-------------------|-------------------------------------|
| Maurice (85) | 2017 | N=481 | Positive surgical margins: more common in MIA vs. OA group (OR 2.0; 95% CI, 1.1–3.6; P=0.03), though the relationship only remained for pT3 disease on subgroup analysis | For ACC confined to the adrenal gland, MIA is comparable to OA with faster post-operative recovery. OA is still superior to MIA regarding locally advanced ACC |
| 320 OA, 161 MIA | | LN dissection was less likely in MIA vs. OA group (OR 0.1, 95% CI, 0.03–0.6; P=0.01), though the relationship was only true in males, tumors ≥10 cm, and cN0 disease | |
| Data obtained from National Cancer Database | | Median LOS was shorter in MIA vs. OA groups (3 vs. 6 d, P<0.01) | |
| Stages I–IV | | Readmissions were non-significantly less in MIA vs. OA groups (4.4% vs. 8.8%, P=0.08) | No differences were found between MIA and OA regarding LN yield or OS |
| Fossà (87) 2013 | N=32 | Smaller tumors in LA vs. OA group | LA provides similar long-term outcomes to OA in addition to significantly superior short-term outcomes (e.g., complication rate, hospital stay). Given that many patients have early relapse regardless of laparoscopic vs. open approach, the short-term benefits should factor into decision-making. |
| 15 OA, 17 LA | | Shorter median operative time (150 vs. 230 min), lower median EBL (400 vs. 1,700 mL), lower median number of perioperative transfusions (0 vs. 3), fewer postoperative complications, and shorter median hospitalization (6 vs. 13 d) after LA vs. OA | |
| Single center | | R0 resection rate: 12/15 (80%) after OA, 12/17 (71%) after LA | |
| Stages I–III | | Recurrence rate: 15/17 (88%) after OA, 12/15 (80%) after LA | Differences in median PFS (15.2 months after LA vs. 8.1 months after OA; P=0.057) and median OS (103.6 months after LA vs. 36.5 months after OA; P=0.22) not statistically significant |
| Porpiglia 2010 N=43 (88) | | Recurrence rate: 50% LA vs. 64% OA | OA and LA “may be comparable in terms of recurrence-free survival,” though the data is not conclusive. |
| 25 OA, 18 LA | | Median RFS: 18 months OA vs. 23 months LA (P=0.8) | |
| Surgery performed at multiple centers | | Mortality during follow-up: 28% OA vs. 5% LA | |
| Stages I–II | | Survival time: no differences between groups | |

Table 3 (continued)
Positive surgical margins were more common in the minimally invasive versus open group (OR 2.0; 95% CI, 1.1–3.6; \(P=0.03\)); subgroup analyses demonstrated this was due to the patients with pT3 or greater disease. No statistically significant differences in 3-year OS were found between MIA and OA (58.0% versus 62.1%, \(P=0.42\)) on univariable analysis; adjusted analysis yielded a \(P\) value of 0.05, which did not meet the authors’ pre-set criteria for significance of \(P<0.05\). The authors concluded that MIA provides acceptable long-term outcomes with faster postoperative recovery for patients with Stage I–II ACC.

Finally, Lee and colleagues retrospectively examined 201 patients from multiple centers (n=154 for OA and n=47 for MIA) (83). Importantly, ENSAT staging was not incorporated into this study. This study found no difference in 30-day mortality between the MIA group (29.3%) versus the OA group (30.9%), \(P=0.839\). Intraoperative tumor rupture did not occur more frequently in the minimally invasive versus open groups (12.2% versus 9.4%, \(P=0.612\)) and R0 status was achieved in a comparable number of patients (77.0% after MIA versus 72% after OA), \(P=0.50\). Estimated blood loss (HR 1.013, 95% CI, 1.001–1.026, \(P=0.038\)) and T stage (T3–T4 versus T1–T2: HR 2.102, 95% CI, 1.106–4.348, \(P=0.045\)) were predictive of survival, whereas surgical approach was not (HR for OA versus MIA 2.152, 95% CI, 0.601–7.714, \(P=0.239\)) on multivariable analysis. The authors concluded that MIA can be considered in patients with localized tumors less than or equal to 10 cm.

**Discussion**

Current ESES and ENSAT guidelines strongly recommend OA as the standard of care surgical approach for patients with suspected ACC (19). While there is cautious acknowledgement of LA as an option in carefully selected patients with suspected ACC, the panel recommends against LA for all cases of stage III or stage IV ACC. For suspected ACC less than 6 cm and confined to the adrenal gland (stage I or II), the panel states that LA is an option, but should be undertaken only in high-volume centers. Notably, the panel agreement was low, the strength of the recommendation was weak, and the evidence level supporting LA was rated as very low. Lastly, the necessity of respecting the principles of oncological surgery is emphasized with a low threshold for open conversion if a risk of tumor spillage or capsular disruption is encountered. Notably, the National Comprehensive Cancer Network also recommends OA even for localized ACC, citing "increased
risk for local recurrence and peritoneal spread when done laparoscopically" (21).

When evaluating data from retrospective analyses, it is important to consider their results in the context of any limitations inherent to their study designs. The lack of randomization allows for between-group differences. Indeed, many of the studies comparing OA to LA report significant differences in factors such as patient age, tumor size, and stage at diagnosis. Since they are not controlled nor randomized trials, retrospective studies are prone to confounding errors, in which factors other than the intervention in question affect the outcome under investigation.

Because ACC is exceedingly rare, and because it is only definitively diagnosed on post-resection or autopsy histology, studies evaluating treatment options for ACC are exclusively retrospective. Although all retrospective studies are subject to the errors and biases listed above, each does not control for potential sources of unintended influence with equal success. Furthermore, surface-level consideration of studies published on open versus laparoscopic outcomes for ACC appear to show a similar number of studies landing on either side of the debate; critical analysis of the quality of these studies is therefore paramount in determining which data best approximates an unbiased truth.

Despite any degree of control that some studies attempted to employ, many had results whose generalizability was limited by a small or unbalanced patient population or limited long-term follow-up (24,73,75,77,78,81,84,87,88), a fact acknowledged by the current ESES and ENSAT guidelines (19). In ten studies (24,70,75,76,78,80,82,87-89), 20 or fewer patients were included in one or both of the surgical groups and seven studies included fewer than 50 patients total (24,78-80,82,87,88). Considering the small number of total patients studied, meta-analyses are one way to bolster otherwise equivocal data. Autorino et al. were able to pool results from 9 studies that directly compared LA (240 cases) to OA (557 cases), with each study including a minimum of 10 cases performed via each approach (86). Selection biases that plagued individual studies additionally manifested in this pooled analysis, with significantly smaller tumors seen in the LA group versus the OA group (weight mean difference -3.41 cm, P<0.001). Although no differences in recurrence rates, time to recurrence, or cancer specific mortality were seen between groups, patients who underwent LA did have significantly higher rates of peritoneal carcinomatosis. The finding that increased rates of peritoneal carcinomatosis after LA persisted despite ACCs being significantly larger in the OA group lends credibility to the likelihood that this is a true association.

Studies that included larger numbers of patients were also subject to confounding. Maurice et al. examined data on 481 patients from the National Cancer Database, the largest sample size of all studies comparing LA (n=161) and OA (n=320) for ACC (85). The authors concluded that MIA was appropriate for ACC confined to the adrenal gland. However, statistically significant differences were present in baseline demographics and tumor characteristics including age, Charlson Comorbidity Index, tumor size, T-stage, and whether or not the surgery was performed at a high-volume center; clinical N-stage approached significance in favor of MIA (P=0.08). Though the authors found no difference in 3-year OS between patients undergoing MIA (58.0%) versus OA (62.1%) with P=0.42, this difference favoring OA trended towards statistical significance on adjusted analysis (P=0.05).

An important finding from Miller et al. also calls into question the true accuracy of data collected on outcomes of ACC patients (77). In this study, 13 of 40 patients (30%) with ACC considered pre-operatively to be stage II were upstaged postoperatively to stage III after undergoing LA; this trend held true for patients undergoing OA, with 22 of 71 (31%) pre-operative stage II patients upstaged to stage III ACC postoperatively. Importantly, in these upstaged patients, 75% of patients in the LA group had positive margins versus only 36% in the OA group, leading the authors to state that more complete resection is achieved with an open versus laparoscopic approach. The implication of these data is important because any study concluding that a threshold of stage II disease or no local invasion should be used in determining when to employ LA (24,81,83-85) will have to reconcile the fact that 30% of these tumors may in fact be stage III at surgical resection.

In addition, volume-outcome relationships seen in other major cancer surgeries have been suggested to influence adrenalectomy for suspected ACC (90). One study stratified patients by both procedure type (OA versus LA) and location of surgery (referring hospital versus index hospital) found significant differences between the same type of surgery (OA) performed at a referring hospital versus the index hospital. This included important measures such as R0 margin status (44.3% after OA at referring hospital versus 89.1% after OA at index hospital) and peritoneal carcinomatosis (27.6% after OA at referring hospital versus 19.6% after OA at index hospital) (73). Therefore, where these surgeries are done may greatly influence patient outcomes.

Given that ACC is uncommon and rarely diagnosed pre-
operatively, it is unlikely that a randomized trial of OA versus LA for suspected ACC will ever be conducted. Therefore, we are limited to well-controlled observational studies on which to make treatment recommendations. The priority of any cancer surgery is oncologic control. Given the aggressive nature, poor prognosis, and lack of effective systemic treatment options for ACC, complete surgical resection is the best chance to cure these patients. There is a signal that LA may result in greater likelihood of a positive surgical margin and an increased risk of peritoneal carcinomatosis. Both of these events are uniformly fatal. Therefore, we should not allow surgical hubris to overshadow the underlying aggressive biology of ACC and potentially compromise outcomes for these patients: simply because we can perform LA for ACC does not necessarily mean that we should.

Conclusions

A prospective, randomized clinical trial comparing LA to OA for ACC is unlikely to be performed. Therefore, well-controlled observational studies remain the highest level of evidence to support surgical decision-making in patients with suspected ACC. OA is the definitive surgical standard of care for stages III and IV ACC. While data are less clear for clinical Stage I and II ACC, the risk of pathologic upstaging, tumor spillage, and positive surgical margin status cannot be understated since the latter two are uniformly fatal. Therefore, current guidelines across the world recommend OA for all patients with adrenal tumors suspicious for ACC.

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Footnote

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