



AperTO - Archivio Istituzionale Open Access dell'Università di Torino

European Society of Endocrinology Clinical Practice Guidelines on the management of adrenocortical carcinoma in adults, in collaboration with the European Network for the Study of Adrenal Tumors

This is the author's manuscript

Original Citation:

Availability:

This version is available http://hdl.handle.net/2318/1650497

since 2019-04-29T10:06:15Z

Published version:

DOI:10.1530/EJE-18-0608

Terms of use:

Open Access

Anyone can freely access the full text of works made available as "Open Access". Works made available under a Creative Commons license can be used according to the terms and conditions of said license. Use of all other works requires consent of the right holder (author or publisher) if not exempted from copyright protection by the applicable law.

(Article begins on next page)

European Society of Endocrinology Clinical Practice Guidelines on the Management of Adrenocortical Carcinoma in Adults, in collaboration with the European Network for the Study of Adrenal Tumors

Martin Fassnacht^{1,2*}, Olaf M. Dekkers^{3,4,5}, Tobias Else⁶, Eric Baudin^{7,8}, Alfredo Berruti⁹, Ronald R. de Krijger^{10, 11, 12, 13}, Harm R. Haak^{14,15, 16}, Radu Mihai¹⁷, Guillaume Assie^{18, 19}, Massimo Terzolo^{20*}

¹ Dept. of Internal Medicine I, Div. of Endocrinology and Diabetes, University Hospital, University of Würzburg, Würzburg, Germany

² Comprehensive Cancer Center Mainfranken, University of Würzburg, Würzburg, Germany ³ Department of Clinical Epidemiology, Leiden University Medical Centre, Leiden, the

Netherlands

⁴ Department of Clinical Endocrinology and Metabolism, Leiden University Medical Centre, Leiden, the Netherlands

⁵ Department of Clinical Epidemiology, Aarhus University Hospital, Aarhus, Denmark ⁶ Department of Internal Medicine, Division of Metabolism, Endocrinology and Diabetes, University of Michigan, Ann Arbor, MI, USA

⁷ Endocrine Oncology and Nuclear Medicine, Institut Gustave Roussy, Villejuif, France
 ⁸ INSERM UMR 1185, Faculté de Médecine, Le Kremlin-Bicêtre, Université Paris Sud, Paris, France

⁹ Department of Medical and Surgical Specialties, Radiological Sciences, and Public Health, Medical Oncology, University of Brescia at ASST Spedali Civili, Brescia, Italy.

¹⁰ Dept. of Pathology, Erasmus MC University Medical Center, Rotterdam, The Netherlands

¹¹ Dept. of Pathology, University Medical Center Utrecht, Utrecht, The Netherlands

¹² Dept. of Pathology, Reinier de Graaf Hospital, Delft, The Netherlands

¹³ Princess Maxima Center for Pediatric Oncology, Utrecht, The Netherlands

¹⁴ Department of Internal Medicine, Máxima Medical Centre, Eindhoven/Veldhoven, the Netherlands

¹⁵ Maastricht University, CAPHRI School for Public Health and Primary Care, Ageing and Long-Term Care, Maastricht, the Netherlands

¹⁶ Department of Internal Medicine, Division of General Internal Medicine, Maastricht University Medical Centre+, Maastricht, the Netherlands.

¹⁷ Department of Endocrine Surgery, Churchill Cancer Centre, Oxford University Hospitals NHS Foundation Trust, Oxford, UK

¹⁸ Department of Endocrinology, Reference Center for Rare Adrenal Diseases, Reference Center dor Rare Adrenal Cancers, Hôpital Cochin, Assistance Publique Hôpitaux de Paris, France

¹⁹ Institut Cochin, Institut National de la Santé et de la Recherche Médicale U1016, Centre national de la recherche scientifique UMR8104, Université Paris Descartes, Sorbonne Paris Cité, Paris, France

²⁰ Internal Medicine, San Luigi Hospital, Dept. of Clinical and Biological Sciences, University of Turin, Orbassano, Italy

*corresponding authors

Correspondence should be addressed to Martin Fassnacht (Email fassnacht_m@ukw.de) and Massimo Terzolo (Email: terzolo@usa.net)

1 Abstract

2

3 Adrenocortical carcinoma (ACC) is a rare and in most cases steroid hormone producing 4 tumor with variable prognosis. The purpose of these guidelines is to provide clinicians with 5 best possible evidence-based recommendations for clinical management of patients with 6 ACC based on the GRADE (Grading of Recommendations Assessment, Development and 7 Evaluation) system. We predefined four main clinical questions, which we judged as particularly important for the management of ACC patients and performed systematic 8 9 literature searches: (A) What is needed to diagnose an ACC by histopathology? (B) Which 10 are the best prognostic markers in ACC? (C) Is adjuvant therapy able to prevent recurrent 11 disease or reduce mortality after radical resection? (D) What is the best treatment option for 12 macroscopically incompletely resected, recurrent or metastatic disease? Other relevant questions were discussed within the group. SELECTED RECOMMENDATIONS: (i) We 13 recommend that all patients with suspected and proven ACC are discussed in a 14 15 multidisciplinary expert team meeting (ii) We recommend that every patient with (suspected) 16 ACC should undergo careful clinical assessment, detailed endocrine work-up to identify autonomous hormone excess, and adrenal-focused imaging. (iii) We recommend that 17 18 adrenal surgery for (suspected) ACC should be performed only by surgeons experienced in 19 adrenal and oncological surgery aiming at a complete en-bloc resection (including resection 20 of oligo-metastatic disease). (iv) We suggest that all suspected ACC should be reviewed by 21 an expert adrenal pathologist using the Weiss score and providing Ki67 index. (v) We 22 suggest adjuvant mitotane treatment in patients after radical surgery that have a perceived 23 high risk of recurrence (ENSAT stage III, or R1 resection, or Ki67 >10%). (vi) For advanced 24 ACC not amenable to complete surgical resection, local therapeutic measures (e.g. radiation 25 therapy, radiofrequency ablation, chemo-embolization) are of particular value. However, we suggest against the routine use of adrenal surgery in case of widespread metastatic disease. 26 27 In these patients we recommend either mitotane monotherapy or mitotane, etoposide, 28 doxorubicin, and cisplatin depending on prognostic parameters. In selected patients with a 29 good response, surgery may be subsequently considered. (vii) In patients with recurrent disease and a disease-free interval of at least 12 months, in whom a complete 30 31 resection/ablation seems feasible, we recommend surgery or alternatively other local 32 therapies. Furthermore, we offer detailed recommendations about the management of 33 mitotane treatment and other supportive therapies. Finally, we suggest directions for future 34 research.

35

1. Summary of recommendations

36

37 After the review process all Recommendations without Rational will be provided here as38 summary.

39 40

41 2. Adrenocortical Carcinoma – epidemiology, pathogenesis, clinical 42 presentation, and general prognosis

43

44 Epidemiology and pathogenesis

The estimated incidence of adult adrenocortical carcinoma (ACC) is between 0.7 - 2.0 per 45 million per year {Kebebew, 2006 #3;Kerkhofs, 2013 #2}. ACC can occur at any age with a 46 47 peak incidence between 40 and 60 years, and with women being more often affected (55-48 60%). In adults, the vast majority of ACCs are sporadic. Occasionally, however, they occur 49 as part of hereditary syndromes such as Li-Fraumeni syndrome, Lynch syndrome, multiple 50 endocrine neoplasia (MEN) 1 and familial adenomatous polyposis {Berruti, 2012 #20;Petr, 51 2016 #34}. In recent years several multi-center studies have shed light on the pathogenesis 52 of ACC {de Reynies, 2009 #14; Fragoso, 2012 #18; Ronchi, 2013 #324} {Jouinot, 2017 #17}, but 'multi-omic' studies {Assie, 2014 #12; Juhlin, 2015 #19; Zheng, 2016 #16} reveal that only 53 a minority of ACC cases have pathogenic driver mutations. For details on this topic we refer 54 55 to recent reviews {Assie, 2014 #11;Else, 2014 #135;Faillot, 2016 #277}.

56

57 Clinical presentation (Table 1)

58 ACC may present with autonomous adrenal hormone excess or with symptoms caused by 59 an abdominal mass. An increasing number of cases are diagnosed within the group of 60 incidentally discovered adrenal masses (incidentalomas) (≈ 10-15%). However, the likelihood 61 of an adrenal incidentaloma being an ACC is low {Terzolo, 1997 #359;Cawood, 2009 62 #326;Fassnacht, 2016 #46}. About 50-60% of patients with ACC have clinical hormone 63 excess. Hypercortisolism (Cushing's syndrome), or mixed Cushing's and virilizing syndromes are observed in the majority of these patients. Pure androgen excess is less frequent while 64 estrogen or mineralocorticoid excess are very rare {Seccia, 2005 #360;Fassnacht, 2011 65 66 #61;Else, 2014 #135;Berruti, 2014 #35;Kerkhofs, 2015 #78;Fassnacht, 2013 #60}. Non-67 specific symptoms from an abdominal mass include abdominal discomfort (nausea, vomiting, 68 abdominal fullness) or back pain. Classical malignancy-associated symptoms such as weight 69 loss, night sweats, fatigue or fever are rarely present.

70

71 Table 1: Clinical presentation of ACC[#]

72

Autonomous adrenal hormone excess	50-60 %
Hypercortisolism (Cushing's syndrome)*	50-70 %
Androgen excess (virilization) in female patients*	20-30 %
Estrogen excess (feminization) in male patients*	5 %
Mineralocorticoid excess*	2-3 %
Non-specific symptoms from an abdominal mass	30-40 %
Incidentally detected by imaging for other purpose	10-15 %

73 *# number derived from: {Berruti, 2014 #35;Fassnacht, 2009 #56;Johanssen, 2010 #69}, and the*

74 ENSAT ACC registry

- 75 * frequently combined
- 76
- 77 General prognosis

78 The median overall survival of all ACC patients is about 3-4 years. The prognosis is, 79 however, heterogeneous. Complete surgical resection provides the only means of cure. In 80 addition to radical surgery, disease stage, proliferative activity/tumor grade, and cortisol 81 excess are independent prognostic parameters (see also section 4.2. and 5.5.). Five-year 82 survival is 60-80% for tumors confined to the adrenal space, 35-50% for locally advanced 83 disease, and much lower in case of metastatic disease with reported percentages ranging 84 from 0% to 28% {Icard, 2001 #79;Bilimoria, 2008 #80;Sturgeon, 2006 #81;Fassnacht, 2010 85 #57;Fassnacht, 2009 #58;Fassnacht, 2011 #61;Fassnacht, 2012 #28;Kerkhofs, 2015 #78}. 86

- 86
- 87 88

89 **<u>3. Methods</u>**

90

91 3.1. Guideline working group

This guideline was developed by The European Society of Endocrinology (ESE) in 92 93 collaboration with the European Network for the Study of Adrenal Tumours (ENSAT). The 94 chairs of the working group Martin Fassnacht and Massimo Terzolo as well as the 95 methodological expert Olaf Dekkers were appointed by the ESE Clinical Committee. Tobias Else served as representative of The Endocrine Society, USA, and Radu Mihai as 96 97 representative of the European Society of Endocrine Surgeons. The other members were 98 suggested by the chairs and approved by the Clinical Committee of ESE. The 99 multidisciplinary team consisted of the following experts: endocrinologists (Guillaume Assie 100 (France), Olaf Dekkers (The Netherlands), Tobias Else (USA), Martin Fassnacht (Germany), 101 Harm Haak (The Netherlands), Massimo Terzolo (Italy), oncologists (Eric Baudin (France), 102 Alfredo Berruti (Italy), a pathologist Ronald de Krijger (The Netherlands), and an endocrine 103 surgeon Radu Mihai (UK). The working group had three in-person meetings (November 104 2016, September 2017, and March 2018) and communicated by phone and email. 105 Consensus was reached upon discussion; minority positions were taken into account in the 106 rationale behind recommendations. Prior to the process, all participants completed conflict of 107 interest forms.

108

109 3.2 Target group

This guideline was developed for healthcare providers involved in the care of patients with adrenocortical carcinoma *i.e.*, endocrinologists, oncologists, surgeons, radiologists, nuclear medicine physicians, radio-oncologists, pathologists, specialists in general internal medicine, and nurse specialists. However, general practitioners or gynecologists or dermatologists (who are involved in the diagnostic of androgen excess) might also find the guideline useful, as might our patients. In addition, the guideline document can serve as a source document for the preparation of patient information leaflets.

117

118 3.3 Aims

The overall purpose of this guideline is to provide clinicians with practical guidance for the management of patients with adrenocortical carcinoma. In clinical practice, treatment decisions should take into account the recommendations but also the clinical judgment of the treating physician. Recommendations are thus never meant to replace clinical judgment. In some countries not all recommended tests and treatments, or both, might be available. Thus,
the recommendations have certainly be interpreted in the context of available
resources/treatment in the community, in which the patient is being seen.

126 127

128 **3.4 Summary of methods used for guideline development**

129 The methods used have been described in more detail previously {Bollerslev, 2015 #1}. In 130 short, the guideline used GRADE (Grading of Recommendations Assessment, Development 131 and Evaluation) as a methodological base. The first step was to define clinical question(s) 132 (see section 3.5), the second being a systematic literature search (see Section 3.6). After including all relevant articles, we 1), rated the quality of the evidence, and 2) estimated an 133 134 average effect for specific outcomes (if possible). The quality of evidence behind the 135 recommendations is classified as very low (+OOO), low (++OO), moderate (+++O) and 136 strong (++++).

137 For the recommendations we took into account: 1) quality of the evidence, 2) balance of 138 desirable and undesirable outcomes, 3) values and preferences (patient preferences, goals 139 for health, costs, management inconvenience, feasibility of implementation, etc) {Andrews, 140 2013 #137; Andrews, 2013 #138}. The recommendations are worded as recommend (strong 141 recommendation) and suggest (weak recommendation). The meaning of a strong 142 recommendation can be stated as follows: reasonably informed persons (clinicians, 143 politicians and patients) would want the management in accordance with the recommendation. For a weak recommendation, most persons would still act in accordance 144 145 with the guideline, but a substantial number would not {Andrews, 2013 #138}. Formal 146 evidence syntheses were performed and graded only for recommendations addressing our 147 initial four questions. Recommendations based on good practice and experience of the 148 panelists were not graded {Guyatt, 2015 #139}. Recommendations were derived from 149 majority consensus of the guideline development committee, but if at least one member had 150 substantial disagreements, this is acknowledged in the manuscript. If two or more panelists 151 did not agree with a recommendation, this was considered as not consensus. For 152 transparency, all recommendations are accompanied by text explaining why specific 153 recommendations were made.

154

3.5. Clinical question, eligibility criteria and endpoint definition

At the beginning of the guideline development process, the panel agreed on 30 clinical questions in the management of patients with ACC that should be addressed in the guidelines. In a next step, we agreed on four most relevant clinical questions (Table 2), for which a detailed literature search and review was subsequently performed.

160

161 **3.6 Description of search and selection of literature**

162 A literature search of electronic medical databases was performed for all four clinical 163 guestions. As we expected that single publications could contribute to different questions (for 164 example 2 and 4) we decided to perform one overarching search using broad search terms. 165 The search revealed 5988 papers, of which 615 were duplicates. In summary, we included 18 publications for clinical question 1 (diagnostics for ACC), 35 studies for clinical question 2 166 (prognosis), 10 publications for clinical question 3 (adjuvant therapy) and 48 publications for 167 168 clinical question 4 (recurrent/advanced disease). The review of hormonal overproduction as 169 prognostic factor was published as stand-alone paper {Vanbrabant, 2018 #140}. For question 170 3, we included one study after having been provided with baseline characteristics and adjusted estimates for mitotane therapy not reported in the original publication {Bertherat,2007 #82}.

- 173
- 174

175 **<u>3.7. Review process and endorsement of other societies</u>**

176 A draft of the guideline was reviewed by four experts in the field (see "Acknowledgment" section) and has been submitted for comments by ESE and ENSAT members. In addition, 177 178 the following societies and networks were asked for review and finally endorsed the guidelines: the European Society of Endocrine Surgeons, the Endocrine Society, USA, the 179 180 European Society of Pathology, the American-Australian-Asian Adrenal Alliance (A5), the 181 European Reference Network on Rare Endocrine Conditions (Endo-ERN), the European Reference Network on Rare Adult Solid Cancers (ERN EURACAN). Furthermore, patient 182 groups were approached to review the guidelines. All comments and suggestions were then 183 184 discussed and implemented as appropriate by the panel (all comments and responses are 185 provided in Appendix 8).

187Table 2: Overview of the key clinical questions and predefined outcome parameters

Clinical Question	Predefined selection criteria and key outcome	Metrics of the	
	parameters	literature search	
Question 1:	Population Adrenal masses	Number of papers included:	
Pathology - what is needed to diagnose an ACC?	Restriction Minimum 25 ACC patients	1a: n=4	
Sub-question 1A: How to make a distinction between	• Each marker has to be reported in at least 2 independent cohorts Outcome	1b: n=15 (2 papers contributed to	
adrenocortical/non-adrenocortical tumor?	Diagnostic accuracy (Sensitivity/specificity/NPV/PPV) Diagnostic marker:	both)	
Sub-question 1B How to make a distinction between benign or malignant or indeterminate behavior in adrenocortical tumors	 (Weiss Score), Ki67, reticulin, Helsinki, SF-1, melan A, inhibin, calretinin, chromogranin, SRC1 Reference standard: Weiss-Score¹ Recurrence 		
Question 2:	Population (minimum 100 ACC patients):	Number of papers	
Which are the best prognostic markers in ACC?	 Patients after radically resected ACC Patients with advanced ACC Restriction: 	included: 35	
	Prognostic marker has to be reported in at least 2 independent cohorts		
	Prognostic markers to be considered:		
	Tumor stage (different systems: Sullivan, Lee, UICC, ENSAT, etc.), sex, age, Ki67, hormone section, Weiss score, mitotic index, R status, molecular/immunohistological markers		
	Outcome		
	Overall survival, disease-free and progression-free survival, prognostic ability		
Question 3:	Population:	Number of papers	
	• Diagnosis of ACC with macroscopic radical resection (R0, R1,	included:	
Is adjuvant therapy able to prevent recurrent disease or reduce mortality	Rx) Restriction:	Mitotane n=6	
after radical resection?	 Studies with > 10 patients in the intervention group Only studies providing baseling data per tractment group, and 	Radiation therapy n=4	
	Only studies providing baseline data per treatment group, and providing age and stage adjusted estimates		
	In case of >25% overlap only inclusion of the largest study		

Question 4: What is the best treatment option for macroscopically incompletely resected, recurrent or metastatic disease?	 Intervention: Adjuvant treatment with either mitotane, radiation therapy or cytotoxic chemotherapy Control group: Without therapy or other treatment Outcomes: Disease-free survival, overall survival, quality of life, adverse events Population: Macroscopically incompletely resected, recurrent or metastatic ACC Restriction: Studies > 10 patients in the intervention group. Only studies providing baseline data per treatment group Interventions Cytotoxic drugs including mitotane, surgery, radiation therapy, radiofrequency ablation, chemoembolization Control Not mandatory (single arm cohort studies eligible) Outcome Overall survival, progression-free survival, tumor response, quality of life, adverse events 	Number of papers included: cytotoxic drugs including mitotane: n=27 surgery: n= 16 radiation therapy: n=1 radiofrequency ablation: n=1 radionuclide therapy: n=1
--	--	--

- NPV negative predictive value, PPV positive predictive value, SF-1 steroidogenic factor 1, SRC1 steroid receptor coactivator 1, R status Resection status, R0
- microscopically complete resection, R1 microscopically incomplete resection, Rx uncertain resection status ¹ we are aware that the Weiss score was never properly validated, but we decided that there is no other "gold standard")

193 **<u>4. Summary and conclusions from systematic literature reviews</u>**

194

195 **4.1. Clinical question 1: Pathology**

196 We included 17 publications {Blanes, 2007 #141;Creemers, 2016 #142;Erickson, 2001 197 #143:Arola. 2000 #144;Aubert, 2002 #108:Busam. 1998 #146;Kamio, 1990 198 #148;Komminoth, 1995 #149;Pan, 2005 #150;Rubin, 2016 #151;Sbiera, 2010 199 #15;Stojadinovic, 2003 #152;Volante, 2006 #153;Wajchenberg, 2000 #154;Wang, 2014 200 #155;Zhang, 2008 #156;Kovach, 2015 #353} that contributed data to either the diagnosis of 201 ACC in the context of adrenal vs. non-adrenal distinction (4 studies), or in the context of 202 benign vs. malignant adrenocortical tumor distinction (15 studies) (two of them contributing to 203 both subquestions {Arola, 2000 #144;Pan, 2005 #150}). Details of studies are shown in 204 Appendix 1 (in all samples, diagnosis based on histological examination). Melan-A and 205 inhibin-alpha were studied in three publications; all other markers were studied in one or 2 206 publications only. In total data for twenty-seven diagnostic markers were reported. Since 207 many publications included patients who did not reflect the target population in question for 208 this guideline (i.e. patients with a suspicion for ACC), positive or negative predictive values 209 were not provided. A formal meta-analysis was not performed given the low number of 210 studies per marker. Importantly, no study reported on the combined diagnostic ability of a set 211 of markers, which actually may reflect the approach in clinical practice.

212

213 **4.2. Clinical question 2: Prognostic factors**

214 Thirty-five studies reporting on risk factors for recurrence and/or mortality, and that included more than 100 patients with histologically proven ACC, were analyzed {Amini, 2016 215 #157;Asare, 2014 #158;Assie, 2007 #114;Ayala-Ramirez, 2013 #160;Berruti, 2014 216 #35;Beuschlein, 2015 #50;Bilimoria, 2008 #80;Canter, 2013 #164;Duregon, 2017 217 218 #110;Erdogan, 2013 #55;Ettaieb, 2016 #117;Fassnacht, 2009 #58;Freire, 2013 #169;Gicquel, 2001 #170;Glover, 2015 #171;Gonzalez, 2007 #172;Icard, 2001 #79;Jouinot, 219 2017 #17;Kebebew, 2006 #3;Kendrick, 2002 #176;Kim, 2016 #177;Kim, 2017 #178;Libe, 220 221 2015 #29;Livhits, 2014 #180;Lucon, 2002 #181;Margonis, 2016 #182;Margonis, 2016 #183;Millis, 2015 #184;Paton, 2006 #185;Pennanen, 2015 #109;Schulick, 1999 #187;Tran, 222 223 2016 #188;Xiao, 2015 #189;Zini, 2009 #190;Ronchi, 2012 #321}(see Appendix 2 for details 224 of studies included, and Appendix 3 for an overview of all prognostic factors studied). The 225 threshold of 100 cases was defined upfront as with n=100 and an expected number of 226 deaths of 50, statistical power was considered sufficient. Almost all studies reported age, sex 227 and tumor stage as prognostic factors, although several different staging systems were used. 228 A formal comparison of the studies was difficult due to heterogeneity regarding clinical 229 characteristics, use of varying definitions of characteristics (e.g. stage) and different cut-offs 230 (e.g. tumor size, age). Furthermore, the multivariable models presented include adjustment 231 for different additional variables. We acknowledge a concern over the number of variables 232 included in models relative to the number of events, and that this may have the potential to 233 lead to false positive results.

The association between staging and prognosis was robust (+++O), despite different systems being used {Macfarlane, 1958 #281;Sullivan, 1978 #282;Lee, 1995 #283;DeLellis, 2004 #284;Asare, 2014 #158;Miller, 2010 #90;Lughezzani, 2010 #92;Fassnacht, 2009 #58;Libe, 2015 #29;Lam, 2017 #285}. In a formal comparison, the ENSAT staging {Fassnacht, 2009 #58} was slightly superior to the UICC staging {Lughezzani, 2010 #92}. Additionally, the association between hypercortisolism and mortality was consistent, and remained with a positive hazard ratio after adjustments for tumor stage HR 1.71, 95% CI 1.18-2.47 {Vanbrabant, 2018 #140}. Ki67 was studied in five publications, showing worse
prognosis with increasing Ki67 in all studies. Other molecular markers have been studied in
single cohorts only (Appendix 2+3).

It is important to mention that relative risks, even if statistically significant, cannot inform clinical decision making unless translated into predictive values or incorporated in prediction models. Only one study presented a formal prediction model (including the variables tumor size, stage, mitotic index, venous invasion, and endocrine activity), showing a sensitivity of 0.91 and a specificity of 0.90 {Freire, 2013 #169} Another study provided nomograms to facilitate prognosis in individual patients {Kim, 2016 #177}. None of these models, however, has been validated externally.

251

4.3. Clinical question 3: Adjuvant therapy

No randomized clinical trial has been published yet exploring adjuvant therapies; no studies 253 254 comparing quality of life after different treatment modalities were found. We included six 255 studies that assessed the effect of mitotane on recurrence and mortality (Berruti, 2017 256 #82;Beuschlein, 2015 #50;Else, #22;Bertherat, 2007 2014 #125;Grubbs, 2010 257 #191;Postlewait, 2016 #192}. See Appendix 4 for details and Appendix 5 for risk of bias 258 assessment. Due to an overlap of the study population of >25% between studies {Berruti, 259 2017 #22;Bertherat, 2007 #82;Beuschlein, 2015 #50} only the German study cohort from 260 Beuschlein et al. was considered, but not the validation cohort {Beuschlein, 2015 #50}. In 261 one study, forty-seven patients were enrolled in 4 Italian centers where adjuvant mitotane 262 was routinely recommended, 55 patients in 4 Italian centers where no adjuvant strategy was 263 undertaken (control group 1), and 75 German patients left untreated after surgery (control 264 group 2) {Berruti, 2017 #22;Terzolo, 2007 #33}. However, only the most recent update of 265 these series was included in the analysis {Berruti, 2017 #22}. In order to avoid counting data 266 twice only control group 1 was included.

267 In a meta-analysis the pooled hazard ratio for recurrence was 0.7, 95%CI 0.5-1.1; for 268 mortality (5 studies) the pooled hazard ratio was 0.7, 95%CI 0.5-0.9 (Figure 1). All six studies 269 were non-randomized with the potential of a (residual) confounding effect, meaning that 270 treatment choices are based on prognosis (such as performance status of the patient, tumor 271 stage etc.), which introduces imbalance in prognostic factors. It is known that when studying 272 therapeutic effects this confounding effect is difficult to remedy statistically {Bosco, 2010 273 #193}. One study {Berruti, 2017 #22} circumvented the confounding effect by comparing two 274 treatment strategies applied in different settings; such comparison relies on other 275 assumptions {Hernan, 2006 #354}. A further bias in this context is immortal time bias, which can occur if treatment is initiated after follow-up time starts and this is not accounted for in 276 the analysis. Such biases tend to overestimate treatment effects {Suissa, 2008 #194}, and 277 278 were not explicitly accounted for in most studies. Only one study applied a landmark analysis 279 to address this bias {Berruti, 2017 #22}. The overall quality rating was very low (+OOO).

280 Four studies assessed the impact of adjuvant radiation therapy {Fassnacht, 2006 281 #126;Habra, 2012 #123;Else, 2014 #125;Sabolch, 2015 #127}. See Appendix 4 for details 282 and Appendix 5 for risk of bias assessment. The study by Sabolch et al. {Sabolch, 2015 #127} was only considered for data on local recurrence, not for recurrence and mortality 283 284 given the overlap with another study of the same group {Else, 2014 #125}. All but one study (59 patients treated with adjuvant radiation therapy {Else, 2014 #125} were small. We found 285 286 a pooled hazard ratio of 0.8 (95% CI 0.6-1.1) for recurrence and for mortality of 1.0 (95% CI 287 0.7-1.5)(Figure 1). The pooled hazard ratio for local recurrence (three studies) after treatment 288 with radiotherapy was 0.3 (93% CI 0.1-1.9).

All studies were observational with the potential of (residual) confounding effects, immortal

290 time bias was not explicitly accounted for in most studies, and the studies were small with

291 imprecise effect estimates; the overall quality rating was therefore very low (+OOO).

Adjuvant % ES (95% CI) Weight Author Therapy Adjustment Year Mitotane 0.3 (0.2, 0.6) 15.42 Berruti 2017 Mitotane Multivariable Bertherat 2007 Mitotane Multivariable 1.3 (0.8, 2.0) 16.80 Beuschlein German cohort 2015 Mitotane 0.7 (0.5, 1.1) 17.43 Multivariable Flse 0.7 (0.5, 1.0) 19.13 2014 Mitotane Multivariable Grubbs 2010 Mitotane Multivariable 0.5 (0.3, 0.9) 14.28 Postlewait 2016 Mitotane Multivariable 1.4 (0.9, 2.2) 16.94 Subtotal (I-squared = 78.3%, p = 0.000) 0.7 (0.5, 1.1) 100.00 Radiotherapy Else 2014 Radiotherapy Multivariable 0.7 (0.5, 1.1) 51.53 Fassnacht 2006 Radiotherapy Matching 0.8 (0.5, 1.3) 34.03 Habra 2013 Radiotherapy Multivariable 1.3 (0.6, 2.7) 14.45 Subtotal (I-squared = 0.0%, p = 0.417) 0.8 (0.6, 1.1) 100.00 NOTE: Weights are from random effects analysis 1 .25 5 1 5 Treatment Better No Treatment Better

Recurrence in the adjuvant setting

Mortality in the adjuvant setting

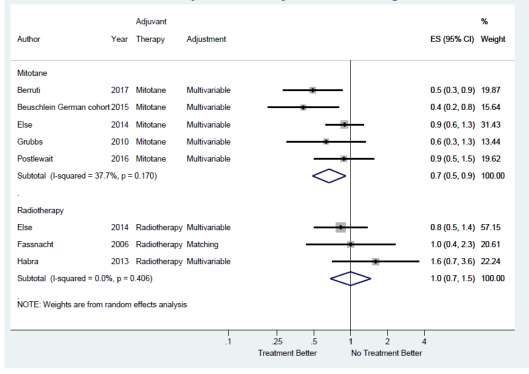


Figure 1 Meta-analysis of recurrence (A) and mortality (B) of included studies on adjuvant therapy after radical resection in ACC

300 **4.4. Question 4: Therapy for advanced or recurrent disease.**

301 A total of twenty-seven publications reported outcomes of 29 different systemic therapies for 302 advanced or recurrent ACC {Berruti, 2005 #24;Fassnacht, 2015 #27;Fassnacht, 2012 303 #28;Gonzalez, 2007 #172;Hermsen, 2011 #68;Sperone, 2010 #32;Abraham, 2002 304 #195;Baudin, 2002 #196;Baudin, 2001 #197;Berruti, 2012 #23;Bonacci, 1998 305 #199;Bukowski, 1993 #200;Decker, 1991 #201;Haak, 1994 #202;Haluska, 2010 #204;Khan, 306 2004 #205;Kroiss, 2016 #206;Kroiss, 2012 #207;Naing, 2013 #208;O'Sullivan, 2014 307 #209;Quinkler, 2008 #74;Schlumberger, 1991 #211;Urup, 2013 #212;Williamson, 2000 308 #213;Wortmann, 2010 #214;Henning, 2017 #215;Lerario, 2014 #216}; two were randomized controlled trials ({Fassnacht, 2015 #27;Fassnacht, 2012 #28}; see Appendix 6 for details of 309 310 studies included). The first randomized trial compared mitotane plus a combination of 311 etoposide, doxorubicin, and cisplatin (EDP-M) to mitotane plus streptozocin in 204 patients 312 with advanced ACC {Fassnacht, 2012 #28}. The trial showed a positive effect of EPD-M on 313 progression-free survival HR 0.55 (95% CI, 0.43 to 0.69; P<0.001), but failed to show a 314 significant effect on mortality (HR 0.79; 95% CI, 0.61 to 1.02; p=0.07); (+++O). The second 315 randomized trial compared linsitinib to placebo (total 139 patients, 2:1 randomization to 316 therapy) and did not show a clear effect on either progression free (HR 0.83, 95% CI 0.56-317 1.21; p=0.30) or overall survival (HR 0.94; 95%CI 0.61-1.44; p=0.77){Fassnacht, 2015 #27}; 318 (+++0).

319 Many publications reported on single arm studies of different therapeutic regimens. These 320 single arm studies have an inherent risk of selection bias, and direct comparison is not 321 possible. Differences in patient characteristics, definition of response criteria and follow-up 322 duration are a concern (+OOO). Given the uncontrolled design a final conclusion about the 323 optimal treatment for advanced recurrent ACC cannot be given. Figure 2 shows response 324 rates from all studies with data for at least one regimen. For most regimens at least some 325 responses (partial or even complete) were reported; treatment merits in case of stable 326 disease is more difficult to judge as this depends highly on duration of follow-up and biology 327 of the disease. Adverse effects from chemotherapy, however, are common and diverse (see 328 Appendix 6).

<u>Study</u>

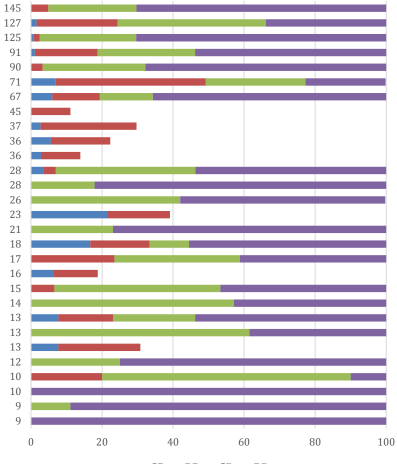
Therapy

Henning, 2017 (Henning, 2017 #215} Fassnacht, 2012 A {Fassnacht, 2012 #28} Fassnacht, 2012 B {Fassnacht, 2012 #28} Hermsen, 2011 {Hermsen, 2011 #68} Fassnacht, 2015 (Fassnacht, 2015 #27} Berruti, 2005 (Berruti, 2005 #24} Gonzalez, 2007 (Gonzalez, 2007 #172} Williamson, 2000 {Williamson, 2000 #213} Bukowski, 1993 (Bukowski, 1993 #200} Decker, 1991 B {Decker, 1991 #201} Abraham, 2002 (Abraham, 2002 #195} Sperone, 2010 (Sperone, 2010 #32} Kroiss, 2012 {Kroiss, 2012 #207} Naing, 2013 {Naing, 2013 #208} Haak, 1994 {Haak, 1994 #202} Kroiss, 2016 (Kroiss, 2016 #206} Bonacci, 1998 (Bonacci, 1998 #199} Urup, 2013 {Urup, 2013 #212} Decker, 1991 A {Decker, 1991 #201}

Gemcitabine and capecitabine
etoposide, doxorubicin, cisplatin, and mitotane
Streptozocin and mitotane
Mitotane and different cytotoxic drug
Linsitinib
Etoposide, doxorubicin, cisplatin, and mitotane
Mitotane
Cisplatin and etoposide
Cisplatin and mitotane
Mitotane
Doxorubicin, etoposide, vincristine, and mitotane
Gemictabine and capecitabine/5-fluorouracil
Sunitinib
Cixutumumab and temsirolimus
Mitotane
Trofosfamide
Etoposide and cisplatin
Cisplatin and docetaxel

Cisplatin and docetaxel

Doxorubicin



■CR ■PR ■SD ■PD

Ν

Lerario, 2014 {Lerario, 2014 #216}	Cixutumumab and mitotane
Haluska, 2010 {Haluska, 2010 #204}	Figitumumab
Schlumberger, 1991 {Schlumberger, 1991 #211}	5-fluorouracil, doxorubicin, and cisplatin
O'Sullivan, 2014 {O'Sullivan, 2014 #209}	Axitinib
Baudin, 2001 {Baudin, 2001 #197}	Mitotane
Baudin, 2002 {Baudin, 2002 #196}	Irinotecan
Kahn, 2004 {Khan, 2004 #205}	Vincristine, teniposide, cisplatin, and cyclophosphamide
Wortmann, 2010 {Wortmann, 2010 #214}	Bevacizumab and capecitabine
Quinkler, 2008 {Quinkler, 2008 #74}	Erlotinib and gemcitabine
Berruti, 2012 {Berruti, 2012 #23}	Sorafenib and metronomic paclitaxel

330 Figure 2: Overview of the objective response rates in studies with systemic therapies in ACC

331 The studies are ordered by number of included patients per regimen. This figure has to be interpreted very cautiously, because study protocols, patient cohorts

332 and characteristics as well as outcome measurements are quite different precluding a direct comparison. CR: complete response; PR: partial response; SD:

333 stable disease; PD: Progression of the Disease. Some of the older studies did not report stable disease or progression, thus these columns don't sum up to 100%

334 Sixteen studies focused on surgery in recurrent and advanced ACC; six publications reported 335 on oligo-metastasectomy (lung, liver) {Datrice, 2012 #118;Gaujoux, 2012 #218;Kemp, 2011 #219;Kwauk, 1993 #220;op den Winkel, 2011 #221;Ripley, 2011 #222}, whereas 10 336 337 publications assessed the effect of surgery in local recurrent and/or metastatic disease 338 {Bellantone, 1997 #223;Crucitti, 1996 #83;Dy, 2013 #225;Erdogan, 2013 #55;Gonzalez, 339 2007 #172; Jensen, 1991 #228; Schulick, 1999 #187; Simon, 2017 #136; Tran, 2013 #231; Dy, 2015 #232}. In patients with metastasectomy 5-survival rates up to 40% were reported 340 {Datrice, 2012 #118;Gaujoux, 2012 #218}, although control groups were lacking (+OOO). 341 342 There were large differences regarding extent of disease, indication, and concurrent 343 treatment in studies comparing a surgical approach to a non-surgical approach for recurrent 344 or advanced disease. The reported benefit of surgery is confounded by differing indications 345 for surgery, and this precludes firm conclusions from being drawn (+OOO). Therefore, the 346 main conclusion is that in patients deemed radically operable by the surgeon/team operation 347 is a treatment option. However, beside prognostic factors like Ki67 a key influencing factor in 348 case of recurrence is the disease-free interval prior to recurrence.

For radionuclide therapy {Hahner, 2012 #373}, transcatheter arterial chemoembolization
{Cazejust, 2010 #233}, radiofrequency ablation {Wood, 2003 #235} and radiation {Ho, 2013
#234} only one small study per procedure was found, and no conclusions can be drawn.

- 352 353
- 354

355 **<u>5. Recommendations</u>**

356

357 **5.1. General remarks**

The main part of this guideline addresses the management of adult patients with ACC. We divided the 62 recommendations in 12 sections. In addition, we provide two flow-charts on the management of patients with ACC amenable to radical resection (Figure 3) and on the management of patients with advanced ACC not amenable to radical resection (Figure 4) to give an efficient overview. However, we would like to emphasize once more that none of these flow-charts nor the entire recommendations can replace clinical judgment of the treating physician and joint decision-making with the patient.

365 366

368

367 **5.1. Overarching recommendations**

369 R.1.1. We recommend that all patients with suspected and proven adrenocortical carcinoma (ACC) are discussed in a multidisciplinary expert team meeting 370 371 (including health care providers experienced in care of adrenal tumors, 372 including at least the following disciplines: endocrinology, oncology, 373 pathology, radiology, surgery) at least at the time of initial diagnosis. In 374 addition, this team should have access to adrenal-specific expertise in 375 interventional radiology, radiation therapy, nuclear medicine, and genetics as 376 well as to palliative care teams.

377378 Reasoning:

379 Despite the lack of studies, the panel was convinced that patients with ACC benefit from 380 multidisciplinary management by a team of experts with experience in care for patients with 381 this rare disease. Ideally, all patients would be managed by such a team throughout the 382 course of their disease, because during the follow-up considerations of multiple diagnostic 383 and treatment modalities might be required. However, in many health care settings this is yet 384 an unrealistic expectation. Therefore, we envision that in the future at least one reference 385 center, that fulfills the above-mentioned criteria, will be established in every country. We 386 believe that it is crucial that every case of suspected ACC is discussed in detail with a panel 387 of experts for this disease at the time of the initial diagnosis. Additionally, this expert team 388 should be ideally requested every time progress is documented (or suspected) and new treatment options might be required. If there is no accessible center with all the required 389 390 expertise in all disciplines, or the patient is not able to travel to such a center, telemedicine 391 approaches should be encouraged to compensate for these limitations.

392 393

R.1.2. We suggest that at any time of decision-making regarding therapy, enrollment in
 a clinical trial (if available) should be considered. Furthermore, we encourage
 patients' participation in registries and the collection of biological material as
 part of structured research programs aimed at defining biomarkers of
 diagnosis, prognosis and treatment response.

399

400 <u>Reasoning:</u>

401 As described above, the evidence for almost all therapeutic strategies for ACC is very low. 402 Furthermore, the efficacy of systemic therapies is limited, including the most commonly used 403 treatments - mitotane and platinum-based chemotherapies, with response rates clearly less 404 than 30% {Baudin, 2001 #197;Berruti, 2012 #20;Else, 2014 #135;Fassnacht, 2012 405 #28;Hermsen, 2011 #68;Hahner, 2005 #64}. Thus, improved treatment paradigms are 406 needed urgently. Clinical trials are the best way to improve our knowledge and patient care. 407 However, the benefits and risk for the individual patient have to be weighed against available 408 data of agents with known or predicted efficacy in ACC.

409 Because of the rarity of the disease, it is crucial to include as many patients as possible in 410 research programs for multicenter therapeutic trials, as well as studies for diagnostic, 411 prognostic and predictive markers. A list of ongoing trials is accessible on https://www.clinicaltrials.gov/. Biological material may include tumor samples, ideally frozen 412 413 and paraffin-embedded, blood-derived and urine samples. National and international 414 research networks such as ENSAT (www.ensat.org){Stell, 2012 #91} and the recently 415 founded A5 (https://adrenal-a5.org/) play instrumental roles in coordinating research 416 programs. Centers providing care to patients with ACC should register as investigators with 417 ongoing trials and also facilitate the collection a of biological material and ensure appropriate 418 consent.

419

420

421

422 **5.2. Diagnostic procedures in suspected ACC**

- 423
- R.2.1. The diagnosis of ACC is not always obvious. We recommend establishing as
 soon as possible whether an adrenal mass is malignant, using all required
 diagnostic tools in a timely fashion.
- 427 428 <u>Reasoning</u>

429 Due to the potentially poor prognosis of ACC, it is critical to know as early as possible if an 430 adrenal mass is malignant or not. Therefore, even if there is only a small likelihood that an 431 adrenal mass is an ACC, this diagnosis should be rapidly excluded with the highest possible 432 certainty. A particular suspicion for an ACC might arise from clinical aspects (e.g. rapidly 433 developed features of adrenocortical hormone excess, see R.2.2), or results from hormonal 434 work-up (see R.2.3), or indeterminate or suspicious imaging (see R.2.4). An adrenal biopsy 435 should only be considered in those selected cases in which an adrenal metastasis of an 436 extra-adrenal malignancy is suspected or when the tumor is considered as inoperable 437 {Fassnacht, 2016 #46} (for details and explanation see R.2.7). The proposed diagnostic 438 work-up is summarized in Table 3.

Table 3: Diagnostic work-up in patients with suspected or proven ACC $\frac{440}{441}$

Hormonal work up	
Glucocorticoid excess	 1mg dexamethasone suppression test or free cortisol in 24-h urine¹ basal ACTH (plasma)²
 Sex steroids and steroid precursors³ 	 DHEA-S 17-OH-progesterone androstenedione testosterone (only in women) 17-beta-estradiol (only in men and postmenopausal women) 11-deoxycortisol
Mineralocorticoid excess	 potassium aldosterone/renin ratio (only in patients with arterial hypertension and/or hypokalemia)
Exclusion of a pheochromocytoma Imaging	 Fractionated metanephrines in 24h urine or free plasma-metanephrines CT or MRI of abdomen and pelvis, Chest CT FDG-PET/CT⁴ Bone or brain imaging (when skeletal or cerebral metastases are suspected)

¹ The 1-mg dexamethasone test is the preferred method to exclude relevant hypercortisolism.

444 However, if overt Cushing syndrome is evident, then cortisol in 24-h urine might be at least as good to 445 quantify the cortisol excess. Alternatively, salivary or serum bedtime cortisol can be used.

446 ² ACTH can be skipped if hypercortisolism is excluded.

³ The most suitable set of precursors and sex hormones has not yet been established and local

- 448 availability might be taken into account.
- ⁴ 449 ⁴ The panel did not agree on the systematic use of FDG-PET/CT (see R.2.4).
- 450 451

442

R.2.2. We recommend that every patient with (suspected) ACC should undergo careful assessment including case history, clinical examination for symptoms and signs of adrenal hormone excess.

455 456 <u>Rea</u>soning

457 All patients should undergo a careful evaluation with detailed history and physical 458 examination. In particular, patients should be evaluated for rapidly developing Cushing's 459 syndrome (which frequently presents not as 'full blown' Cushing, but rather predominantly 460 with muscle weakness, hypokalemia, wasting and constitutional symptoms), and symptoms and signs of a large abdominal mass. Clinical evaluation should additionally focus on 461 462 symptoms and signs of androgen excess, hirsutism or virilization in women or recent onset of 463 gynecomastia in men, because these might be clinical indicators for an androgen- or 464 estrogen-producing ACC, respectively {Fassnacht, 2004 #59;Allolio, 2006 #236;Else, 2014 465 #135;Fassnacht, 2009 #56;Nieman, 2008 #134;Libe, 2007 #237}. Any evidence of cosecretion of different steroids raises the suspicion of an ACC (especially if sex-hormones are 466 467 involved). In contrast, mild, long standing hirsutism is usually not caused by an ACC, but 468 rather due to (among other diagnoses) polycystic ovary syndrome and non-classical 469 congenital adrenal hyperplasia {Legro, 2013 #238}. Primary aldosteronism is rare in ACC 470 and usually accompanied by severe hypokalemia {Funder, 2016 #239}. However, 471 hypokalemia in ACC is more frequently caused by massive cortisol excess overwhelming the 472 renal 11-β hydroxysteroid dehydrogenase type 2 system.

474

R.2.3. We recommend that all patients with suspected ACC undergo a detailed hormonal work-up to identify potential autonomous excess of glucocorticoids, sex-hormones, mineralocorticoids and adrenocortical steroid hormone precursors (see Table 3). In addition, a pheochromocytoma must be excluded.

479 480 Reasoning

481 A comprehensive endocrine work-up is helpful for various reasons. (i) The diagnosis of 482 steroid excess is frequently able to establish the adrenocortical origin of the tumor. (ii) The 483 steroid pattern may indicate whether an adrenal lesion is an ACC. For example, autonomous 484 co-secretion of androgens and cortisol in any patient and secretion of steroid precursors or 485 estradiol in males are highly suspicious for ACC {Fassnacht, 2004 #59}. Furthermore, 486 hormonal evaluation is of prognostic value as cortisol-secreting tumors generally have a 487 worse prognosis {Vanbrabant, 2018 #140}. (iii) If undiagnosed, autonomous cortisol 488 secretion may be followed by life-threatening adrenal insufficiency after complete resection of 489 the primary tumor. The best test to diagnose autonomous cortisol secretion is the 1-mg 490 overnight dexamethasone suppression test {Nieman, 2008 #134}. If hypercortisolism is 491 present, it is crucial to prove ACTH-independency, because an adrenal metastasis of an 492 ectopic ACTH-secreting tumor (e.g. lung cancer) can mimic an ACC. (iv) Elevated hormones 493 prior to surgery may serve as tumor markers during follow-up. Finally, conventional imaging 494 cannot discriminate an ACC from a pheochromocytoma. However, undiagnosed 495 pheochromocytoma may lead to dangerous hypertensive crises (especially during invasive 496 procedures). Therefore, a pheochromocytoma has to be ruled out in any case of an adrenal 497 tumor whenever no obvious autonomous steroid excess is present {Fassnacht, 2016 #46}. It 498 is important to note, however, that slightly elevated metanephrines levels (< 2-fold), 499 particularly when inconsistent with a large tumor size, might be non-specific and can be 500 observed in ACC.

501

502

503 R.2.4. We recommend adrenal-focused imaging in all patients with suspected ACC.

504

505 Reasoning

506 Imaging tools for adrenal tumors were carefully reviewed during the development of the ESE-507 ENSAT guidelines for adrenal incidentalomas {Dinnes, 2016 #54;Fassnacht, 2016 #46}. 508 Thus, we refer to these documents for details. Briefly, there are currently three main imaging 509 techniques available for the differentiation of malignant and benign adrenal tumors: 510 computed tomography (CT), magnetic resonance imaging (MRI), and positron emission 511 tomography with ¹⁸F-2-deoxy-D-glucose (mostly combined with CT; FDG-PET/CT). CT and 512 MRI are techniques mainly optimised to identify benign lesions, providing a tool for the exclusion of adrenal malignancy {Peppercorn, 1998 #244;Caoili, 2002 #243;Blake, 2006 513 #242;Ilias, 2007 #245}. Conversely, FDG-PET/CT is mainly used for the detection of 514 515 malignant disease {Mackie, 2006 #121;Groussin, 2009 #247;Deandreis, 2011 #246}. A 516 recently performed meta-analysis indicated that the level of evidence is low to very low for all 517 these imaging methods {Dinnes, 2016 #54}. In the last 2 years some additional studies have been published {Cistaro, 2015 #362;Altinmakas, 2017 #254;Ciftci, 2017 #255;Bluemel, 2017 518 519 #250;Werner, 2016 #248;Wu, 2016 #249;Nakajo, 2017 #257;Guerin, 2017 #256;Marty, 2018 520 #251;Kim, 2018 #252;Delivanis, 2018 #253;Romeo, 2018 #258;Thomas, 2018 #259;Ng, 521 2018 #260;Kim, 2018 #252}. However, the panel still considers that of the available imaging

522 modalities, only non-contrast CT is sufficiently reliable to rule-out an ACC when the adrenal 523 lesion is homogenous and has low CT density \leq 10 HU. In contrast, ACCs are usually large 524 and of inhomogeneous appearance, and characterized by low fat content (and hence higher 525 HU density){Petersenn, 2015 #323}. Recently, FDG-PET has been proposed as possibly the 526 best second-line test to assess indeterminate masses by unenhanced CT {Cistaro, 2015 527 #362;Guerin, 2017 #256;Nakajo, 2017 #257}. However, the experience shows that sensitivity 528 and negative predictive value are much better than specificity or positive predictive value. 529 Therefore, no consensus could be reached for a general recommendation on FDG-PET in all 530 patients. Additional reasons in favor of systematic FDG-PET are: whole body imaging 531 (beyond thorax and abdomen, particularly for distant bone metastasis) and in advanced 532 disease, a reference uptake value for all metastases can be established, which can help 533 judging the future evolution of disease. Evidence against FDG-PET includes cost, additional 534 radiation exposure, false-positive findings, and difficult access in some countries.

535 If adrenal imaging indicates an indeterminate mass, other parameters should be considered: 536 For instance, in such a situation a tumor size > 4 cm, combined adrenocortical hormone 537 excess (see also R.2.3), rapidly developing symptoms or young age (e.g. < 40 years) might 538 point to an ACC. However, it is important to note that no single imaging method can 539 definitively prove the diagnosis of ACC.

540 541

542R.2.5. We recommend in any case where there is high suspicion for ACC performing a543chest CT, in addition to an abdominal-pelvic cross-sectional imaging (CT or544MRI), because the results might influence therapeutic decision-making.

545

546 <u>Reasoning:</u>

547 Since decisions for treatment strategy, particularly decisions for surgery, and prognostication 548 rely on tumor stage, it is mandatory to systematically and rapidly evaluate for metastases, 549 before initiation of any anti-tumor treatment. Thoraco-abdomino-pelvic imaging will cover the 550 vast majority of metastatic locations, which most often are lung and liver, and will assess 551 locoregional tumor extent. Imaging should include contrast-enhanced imaging. For 552 abdominal imaging there are advantages and disadvantages for both CT and MRI, but for 553 thoracic imaging CT is the method of choice, because it outperforms all other methods in 554 detecting small pulmonary lesions.

- 555 Additional imaging may be required to better characterize tumor vascularization, or specific 556 tumor extent such as a vena cava thrombus.
- 557 558

561

559R.2.6. We suggest performing additional imaging (e.g. bone and brain imaging) only in560case of clinical suspicion of metastatic lesions.

562 <u>Reasoning:</u>

563 Bone and brain metastases are rare events (especially in patients without other metastatic 564 lesions). Therefore, additional imaging focusing on these sites is only warranted when there 565 is increased clinical suspicion or other imaging is suggestive for bone metastases. It should 566 be noted, however, that the basis for this advice has never been studied systematically.

567

R.2.7. We recommend against the use of an adrenal biopsy in the diagnostic work-up of patients with suspected ACC unless there is evidence of metastatic disease that precludes surgery and histopathologic proof is required to inform oncological management.

573

574 <u>Reasoning:</u>

575 Differentiating benign from malignant adrenocortical tumors is very challenging on a biopsy only and may lead to misdiagnosis {Bancos, 2016 #49;Fassnacht, 2016 #46}. Furthermore, 576 577 the biopsy comes with significant risks such as hemorrhage {Williams, 2014 #262}. The risk 578 of tumor dissemination precluding a R0 resection is very low {Williams, 2014 #262}. 579 However, a biopsy might be indicated in an adrenal mass without any hormone excess in 580 patients with a history of extra-adrenal cancers to exclude or prove an adrenal metastasis of 581 an extra-adrenal malignancy. For details see the adrenal incidentaloma guidelines 582 {Fassnacht, 2016 #46}.

- 583
- 584 585

586 **5.3. Surgery for suspected localized ACC**

587

R.3.1. We recommend that adrenal surgery for suspected/confirmed ACC should be performed only by surgeons experienced in adrenal and oncological surgery.

591 <u>Reasoning</u>

ACC surgery requires expertise in both adrenal and oncological surgery due to the specific anatomy, the malignant character of the disease and the potential need for multi-organ enbloc resection to optimize the probability of a R0 resection and minimize the risk of complications.

596 Data comparing outcome between 'high-volume' and 'low-volume' centers for ACC are 597 limited. Published reports from the UK, USA and Spain show an unacceptable low annual 598 workload for the majority of surgeons involved in any adrenal surgery, with a median 1 599 case/year {Palazzo, 2016 #7;Park, 2009 #9;Lindeman, 2018 #264;Villar, 2010 #263}. This 500 situation is likely to have a negative impact on patient care and contrasts significantly with 501 the current status in other surgical specialties.

602 Based on the upper quartile distribution of workload of surgeons in the USA, a volume of 4 603 adrenalectomies/year was used to define a 'high-volume' surgeon {Park, 2009 #9} but this 604 threshold might be too low to inspire confidence. Several studies showed that those doing 605 more than 6-7 cases per year have shorter length of stay and fewer complications {Palazzo, 606 2016 #7;Park, 2009 #9;Gallagher, 2007 #265}. Despite the perceived benefit of being 607 operated in a high-volume center, published data from Italy and the USA showed no 608 significant association between overall survival / disease-free survival and workload even 609 though patients operated in high-volume centers had more radical surgery, more lymph node 610 assessment and more use of chemotherapy {Lombardi, 2012 #266;Gratian, 2014 #268}. In 611 contrast, the creation of national centers for adrenal surgery in The Netherlands led to 612 significantly improved disease-free survival (1y: 93% vs. 78%, 5y: 63% vs. 42 %) {Hermsen, 613 2012 #36;Kerkhofs, 2013 #41}. Therefore, the panel believes that a minimal annual workload 614 of 6 adrenalectomies/year seems to be required to ensure sufficient experience in adrenal 615 surgery, but > 20 adrenalectomies/year are desirable for those involved in surgery for ACC. 616 Furthermore, due to the complexity of some operations, it is essential to involve surgeons

- 617 with different expertise (e.g. vascular, liver, and cardiac surgeons) for pre-surgical planning 618 and during these complex operations.
- 619 Protocols ensuring referral to regional or national centers should be established and patients 620 should feel empowered to ask about the previous experience of individual surgeons.
- 621
- 622

629

R.3.2. We recommend complete *en bloc* resection of all adrenal tumors suspected to
 be ACC including the peritumoral/periadrenal retroperitoneal fat. We
 recommend against enucleation and partial adrenal resection for suspected
 ACC. If adjacent organs are suspected to be invaded, we recommend *en bloc* resection. However, we suggest against the routine resection of the ipsilateral
 kidney in the absence of direct renal invasion.

630 <u>Reasoning</u>

631 Complete resection is of utmost importance for all ACCs and successful surgery is a 632 prerequisite for cure. As the diagnosis of ACC might only become apparent after histological 633 analysis, it remains imperative for all adrenalectomies (laparoscopic or open) in patients with 634 a reasonable suspicion for ACC to respect the principles of oncological surgery in order to 635 ensure complete resection (R0 status) {Gaujoux, 2017 #87;Gaujoux, 2012 #86}.

To ensure that the pathologist can judge the completeness of surgery, any fragmentation of the tumor has to be avoided. Intraoperative tumour rupture or spillage and R2 resection are associated with very high recurrence rates and poor overall survival {Bilimoria, 2008 #80} {Crucitti, 1996 #83}.

640 Although there are no specific studies comparing outcome of surgery with and without 641 resection of invaded adjacent organs, it is deemed to be 'good surgical practice' to resect 642 adjacent tissues that are/could be invaded by tumor. This holds true for involvement of 643 spleen, distal pancreas, stomach, kidney, right liver, colon, diaphragm, the wall of the IVC or 644 left renal vein. A cohort study compared the oncological results of patients with stage II ACC 645 treated by radical adrenalectomy alone or by en-bloc resection with kidney. The results did 646 not support the hypothesis that nephrectomy improves the oncological outcome {Porpiglia, 647 2016 #31}. Combined nephrectomy, however, offers a lower risk of capsular rupture and can 648 include complete lymphadenectomy of the renal hilum, but impairs kidney function and this 649 may limit further access to chemotherapy. Thus, in case of possible invasion in the kidney, 650 partial nephrectomy should be considered on an individual basis.

651 652

R.3.3. Open surgery is the standard surgical approach for confirmed or highly
 suspected ACC. Therefore, we recommend open surgery for all tumors with
 radiological findings suspicious of malignancy and evidence for local invasion.
 However, for tumors < 6 cm without any evidence of local invasion,
 laparoscopic adrenalectomy (respecting the principles of oncological surgery)
 is reasonable if the surgeon has sufficient experience in these types of surgery.

659 660 Reasoning

There is an ongoing debate if laparoscopic adrenalectomy is an acceptable alternative for adrenal tumors with suspicion of ACC. Based on the systematic review on this topic until July 2014 {Fassnacht, 2016 #46} and an additional literature search until December 2017 {Donatini, 2014 #274;Sgourakis, 2015 #273;Autorino, 2016 #272;Langenhuijsen, 2016 665 #271;Lee, 2017 #6;Zheng, 2018 #269;Mpaili, 2018 #270;Huynh, 2016 #333}, we conclude 666 that the quality of evidence from these observational studies is still very low. The main 667 concerns with all these studies are differences of baseline characteristics between groups. 668 and between important prognostic factors, such as tumor stage or size. The lack of any 669 randomized trial prevents any final conclusions. However, in order to provide guidance for 670 clinicians the panel concurs with two other recent European guidelines {Fassnacht, 2016 671 #46;Gaujoux, 2017 #87} and agrees that all tumors with some radiological evidence of local 672 invasion (including enlarged lymph nodes) should undergo surgery with an open approach. 673 The likelihood of a benign adrenal tumor is higher in the group of adrenal incidentalomas ≤ 6 674 cm, for whom a laparoscopic approach is reasonable. However, this cut-off is arbitrary and 675 the experience of the surgeon is the single most important factor. Furthermore, it is advised 676 to convert to an open procedure when obvious signs of invasion are encountered during 677 laparoscopic surgery that would prevent complete resection. For detailed discussion we refer 678 to the recent recommendations for the surgical management of ACC by ESES and ENSAT 679 {Gaujoux, 2017 #87} and the guidelines on adrenal incidentaloma {Fassnacht, 2016 #46}.

Although retroperitoneoscopic adrenalectomy is gaining popularity, only a small number of surgeons are likely to have completed the learning curve to reach sufficient expertise, which is estimated to be at least 20 cases {Barczynski, 2007 #275;Schreinemakers, 2010 #276}. This is a very significant issue in the context of the overall minimal experience of most surgeons offering adrenalectomy (see above). Outside specialized centers with large volume practice, retroperitoneoscopic adrenalectomy should only be considered for benign tumors <4 cm.

687 688

694

R.3.4. We suggest that routine locoregional lymphadenectomy should be performed with adrenalectomy for highly suspected or proven ACC. It should include (as a minimum) the periadrenal and renal hilum nodes. All suspicious or enlarged lymph nodes identified on preoperative imaging or intraoperatively should be removed.

695 <u>Reasoning</u>

696 Reports from several databases indicated that patients with stage III tumors and positive 697 lymph nodes can have a 10-year overall survival rate of up to 40% after resection 698 {Fassnacht, 2009 #58;Lughezzani, 2010 #92;Libe, 2015 #29;Nilubol, 2016 #8;Saade, 2015 699 #93}. However, the wide range of reported lymph node involvement in ACC (from 4 to 73%) 700 {Icard, 2001 #79:Bilimoria, 2008 #80;Harrison, 1999 #88} demonstrates that regional 701 lymphadenectomy is neither formally performed by all surgeons nor accurately assessed or 702 reported by all pathologists. According to large American and French series, approximately 703 10-30% of patients with ACC had formal lymphadenectomy as part of the tumor resection. 704 reflecting the heterogeneity of operative management {Icard, 2001 #79;Nilubol, 2016 #8}. A 705 minimum of four lymph nodes should be retrieved in order to declare lymph node negative 706 cases {Panjwani, #89} Furthermore, in an analysis of 120 cases identified from a multi-707 institutional database, the benefit of lymphadenectomy on overall survival persisted on 708 multivariable analysis controlling for adverse preoperative and intraoperative factors 709 associated with lymphadenectomy, such as tumor size, palpable mass, irregular tumor 710 edges, suspicious nodes on imaging, and multivisceral resection {Gerry, 2016 #94}. The 711 largest series so far included 283 patients and the resection of more than five lymph nodes 712 reduced also the risk of local recurrence and disease-related death in a multivariate analysis

713 {Reibetanz, 2012 #75}.

However, the panel is not in favor of a repeat surgery if complete adrenalectomy was
performed without lymphadenectomy (e.g. due to perceived benign tumor). The clinical
benefit is uncertain and probably lower than the harm (e.g. delayed adjuvant therapy).

717 718

723

R.3.5. We recommend that individualized treatment decisions are made in cases of tumors with extension into large vessels based on multidisciplinary surgical team. Such tumors should not be regarded 'unresectable' until reviewed in an expert center.

724 <u>Reasoning</u>

725 Extension of ACC into the adrenal vein, renal vein or inferior vena cava occurs in 726 approximately 15-25% {Chiche, 2006 #96;Turbendian, 2010 #95;Fassnacht, 2009 #58}. 727 Venous involvement consists mostly of intravenous tumor thrombus. Thrombectomy might 728 require vena cava cross-clamping above or below the hepatic vein confluence or 729 cardiopulmonary bypass, depending on the upper level of extent of the thrombus. The 730 resection might include a complete thrombectomy, a flush manoeuvre and, occasionally, 731 vascular cuff or prosthetic IVC replacement. A 3-year overall survival rate of about 25% in a large series {Mihai, 2012 #97} encourages the performance of a venous resection in the 732 733 presence of vena cava or renal vein invasion but without distant metastases.

734 735

R.3.6. If the first surgery was suboptimal and macroscopically incomplete (R2 resection), we suggest to discuss repeat surgery in a multidisciplinary expert team.

739 740 Reasoning

There has been no prospective study assessing the benefits (or the lack thereof) of early reoperation in patients whose initial adrenalectomy was incomplete (R2 status). It is the panel's view that such patients should have intensive postoperative monitoring and if local recurrence is detected radiologically, in the absence of other metastases, they should undergo surgery with a curative intent at an expert center, if it is deemed likely to lead to an R0 resection.

747 748

751

R.3.7. We recommend perioperative hydrocortisone replacement in all patients with hypercortisolism that undergo surgery for ACC.

752 <u>Reasoning:</u>

Overt ACTH-independent Cushing's syndrome or biochemical autonomous cortisol secretion might lead to adrenal insufficiency after removal of the adrenal source of cortisol (even in patients with incompletely suppressed ACTH) {Eller-Vainicher, 2010 #4}. Therefore, the group unanimously sees a clear indication of intra- and postoperative glucocorticoid replacement, preferably with hydrocortisone, in all patients with evidence for '(possible) autonomous cortisol secretion' (post-dexamethasone cortisol >50 nmol/L (>1.8 μ g/dL)). This should follow the suggestions for major stress dose replacement as per recent international 760 guidelines {Bornstein, 2016 #314}. Postoperatively, the dose of glucocorticoid should be 761 tapered on an individualized basis by a physician experienced with this clinical scenario. 762 763 764 765 5.4. Pathological work-up 766 767 R.4.1. We recommend that the diagnosis of ACC should be confirmed by 768 histopathology (+++0). 769 770 Reasoning: Histopathology is the gold-standard of diagnosing ACC and should in principle be obtained in 771 772 all patients. For patients deemed operable this will be done on the basis of the resection 773 specimen and for those patients who are inoperable, a biopsy will be taken in accordance 774 with good oncological practice. However, the majority of panelists argued that in selected 775 cases biopsy might be omitted when there is advanced disease with unequivocal ACTH-776 independent cortisol excess, androgen excess (testosterone, DHEAS) or estradiol excess. 777 There is no role for biopsy in a patient who is considered suitable for surgery of the adrenal 778 mass. 779 780 781 R.4.2. We suggest that all adrenal tumors, which cannot be readily classified, and all 782 suspected ACC, should be reviewed by an expert adrenal pathologist (++OO). 783 784 Reasoning: 785 Diagnosing ACC can be challenging and misdiagnoses are relatively frequent events. In 21 786 of 161 of the patients (13%) registered with the German ACC Registry between 2006 and 787 2009, the diagnosis of ACC had to be revised by the reference pathologist {Johanssen, 2010 788 #69}. Similar results were found in a large series from Italy with a rate of misdiagnosis in 26 789 out of 300 cases (9%) {Duregon, 2015 #98}. 790 791 792 R.4.3. We suggest the use of immunohistochemistry for steroidogenic factor-1 (SF1) 793 for the distinction of primary adrenocortical tumors and non-adrenocortical 794 tumors (+000). 795 796 Reasoning: 797 Generally, the distinction between adrenocortical and non-adrenocortical tumors is clear and 798 can be made on the basis of hematoxylin and eosin-stained slides. In case of doubt, on the 799 basis of histology only, whether a tumor originates from the adrenal cortex or not, 800 immunohistochemistry with SF1 is the most sensitive and specific marker currently available 801 to establish if the tumor in question is of adrenocortical origin, with a sensitivity of 98% and a 802 specificity of 100% (Sbiera, 2010 #15). If this marker is not available, we advise a 803 combination of markers, which should include inhibin-alpha, melan-A, and calretinin {Sangoi,

806 807

804

805

following local standard procedures.

2011 #99;Weissferdt, 2014 #100}. Depending on the differential diagnosis, other

immunohistochemistry markers used to make alternative diagnoses may be considered

R.4.4. We recommend the use of the Weiss system, based on a combination of 9 histological criteria that can be applied on hematoxylin and eosin-stained slides, for the distinction of benign and malignant adrenocortical tumors (++OO).

813

808

814 <u>Reasoning:</u>

815 There are many classification systems based on histology and/or a limited number of 816 additional markers for the distinction of benign and malignant adrenocortical tumors. The 817 Weiss system is the most widely used, and although it is not fully standardized {Tissier, 2010 #102;Tissier, 2012 #101} the panel favors use of this score. It should be noted that all 818 819 scoring systems have similar inherent problems. Using the Weiss system, a score of 3 or 820 higher (on a total of 9 criteria, see Table 4) indicates ACC {Weiss, 1984 #104;Weiss, 1989 821 #103}. A score of 2 and 3 may be considered as borderline between benign and malignant 822 tumors (tumors of uncertain malignant potential). In such instance, one of several other 823 classification systems, including the van Slooten index {van Slooten, 1985 #107}, the 824 modified Weiss score {Aubert, 2002 #108}, the Helsinki classification {Pennanen, 2015 825 #109;Duregon, 2017 #110}, and the addition of reticulin stain assessment {Duregon, 2013 826 #26} may be used.

827

Special attention should be paid to histological variants of adrenocortical tumors, mainly
oncocytic tumors, which, because of their specific characteristics, will always have a Weiss
score of least 3, whether they are benign or malignant. For these tumors, an adapted scoring
system should be used, the Lin-Weiss-Bisceglia system {Bisceglia, 2004 #111;Duregon,
2011 #112;Wong, 2011 #113}.

833 Table 4 Histopathologic criteria by Weiss {Weiss, 1984 #104;Weiss, 1989 #103}

The presence of three or more of the following criteria highly correlated with subsequent malignant behavior:

- High nuclear grade (Fuhrman criteria {Fuhrman, 1982 #357})
- > 5 mitoses per 50 high-power field
- Atypical mitotic figures
- < 25% of tumor cells are clear cells
- Diffuse architecture (> 33% of tumor)
- Necrosis
- Venous invasion (smooth muscle in wall)
- Sinusoidal invasion (no smooth muscle in wall)
- Capsular invasion

834

835

836

837R.4.5. We recommend the use of Ki67 immunohistochemistry for every resection838specimen of an adrenocortical tumor (++OO).

Ki67 immunohistochemistry has been proposed for prognostic purposes. Higher Ki67 levels
are consistently associated with poor prognosis. Threshold levels of 10% and 20% have
been considered for discriminating low from high Ki67 labeling index {Beuschlein, 2015
#50;Libe, 2015 #29}. However it is not clear whether any single significant threshold can be
determined (see R.5.2.).

Ki67 labeling has been shown to be unevenly distributed in tumors. Therefore, determination
of the labeling index should be done on whole tumors, with specific attention to the area of
highest Ki67 labeling, preferably by use of an image analysis system {Lu, 2014
#115;Papathomas, 2016 #116}. If only a biopsy is available a low Ki67 labeling may not be
representative and therefore can be misleading.

If Ki67 immunohistochemistry is not available, mitotic count may help in prognostic
stratification of ACC. Mitotic count has been proposed for grading of ACC, using >20 mitoses
per 50 high-power field to define high-grade tumors {Weiss, 1989 #103;Assie, 2007
#114;Miller, 2010 #90}. However, the precise correlation between mitotic count and Ki67
labeling is undetermined.

854 855

R.4.6. We recommend that the pathology report of a suspected ACC should at least contain the following information: Weiss score (including the exact mitotic count), exact Ki67 index, resection status, and pathological tumor stage (indicating invasion or not of the capsule and/or surrounding tissue and solutions) and nodal status (+OOO).

862 Reasoning

The importance of Weiss score and Ki67 index has been discussed in R4.4 and R4.5, respectively. It is important that the exact values are given, because this is of prognostic relevance. Resection status is a major prognostic factor (see R 5.2.). Tumor stage, including nodal involvement, is discussed below (see R.5.1).

867

- 868
- 869
- 870

872

871 **5.5. Staging classification and prognostic factors**

- 873 **R.5.1.** At initial diagnosis, we recommend using the ENSAT staging classification 874 (Table 5) (+++O).
- 875
- 876 <u>Reasoning</u>

877 Tumor staging is the most important prognostic factor. Specifically, the presence of metastases is by far the strongest indicator of poor prognosis. Several staging classifications 878 879 have been proposed {Macfarlane, 1958 #281:Sullivan, 1978 #282:Lee, 1995 #283:DeLellis, 880 2004 #284;Asare, 2014 #158;Miller, 2010 #90;Lughezzani, 2010 #92;Fassnacht, 2009 881 #58;Libe, 2015 #29;Lam, 2017 #285}. Among these, the ENSAT staging classification 882 appears to be the most discriminant, but the differences between staging systems are minor 883 {Fassnacht, 2016 #46}(see also section 4.2.). The panel felt strongly that a one unique 884 staging classification should be adopted across centers in order to improve standardization 885 and documentation of clinical data, and so improve patient care and enhance clinical 886 research.

The ENSAT classification requires extensive imaging prior to surgery (see R.2.4.), systematic lymph node resection, a complete surgical report (see R.3.3 and R.3.4.), and a complete pathological report (see R.4.6.).

- 890
- 891 Table 5: ENSAT staging classification {Fassnacht, 2016 #46}
- 892

ENSAT stage	Definition
	T1, N0, M0
	T2, N0, M0
	T1-T2, N1, M0
	T3-T4, N0-N1, M0
IV	T1-T4, N0-N1, M1

893

894 T1: tumor \leq 5cm; T2: tumor > 5cm; T3: infiltration into surrounding tissue; T4: tumor invasion into 895 adjacent organs or venous tumor thrombus in vena cava or renal vein; N0: no positive lymph node; 896 N1: positive lymph node; M0: no distant metastases; M1: presence of distant metastases.

897 898

R.5.2. At initial diagnosis, we recommend taking the following factors into account when assessing the prognosis and treatment options: tumor stage, resection status, Ki67 index (or mitotic count), autonomous cortisol secretion and the patient's general condition (++OO).

- 903
- 904 <u>Reasoning</u>

Of the many reported prognostic factors tumor stage is the most important, because it reflects tumor extent. Especially the presence of metastases is strongly pejorative (see R.5.1.). Resection status is also a strong prognostic factor {Bilimoria, 2008 #80;Johanssen, 2010 #69;Libe, 2015 #29}, and should be carefully documented in the surgical and pathology reports. Furthermore, several studies have identified Ki67 immunostaining (or mitotic index) as major prognostic factors {Morimoto, 2008 #278;Weiss, 1989 #103;Miller, 2010 #90;Beuschlein, 2015 #50;Libe, 2015 #29}. As revealed by our systemic literature search, hypercortisolism was also one of the most consistent prognostic factors (see section 4.2;
{Abiven, 2006 #279;Berruti, 2014 #35;Vanbrabant, 2018 #140}.

Finally, the patient's general condition is an obvious prognostic factor, especially at advanced
age {Asare, 2014 #158}. It is, however, noticeable that ACC patients often do not show
altered general condition despite advanced disease.

917 From a patient perspective, the panel felt it important to consider two distinct scenarios. First,

918 the risk of recurrence of patients with a localized (stage I-III) disease. For these patients, 919 tumor stage, resection status and Ki67 labeling index are currently the main prognostic 920 factors. This panel proposes to define two classes of localized ACC: low/moderate risk ACC 921 includes stage I-II and R0 and Ki67 \leq 10%, whereas high risk ACC includes stage III, R1, or 922 Ki67 >10%. However, the panel is aware that the dichotomy is arbitrary.

- 923 The second scenario to consider deals with the prognosis of patients with advanced disease 924 (stage IV or recurrent disease not amenable to complete resection or R2 resection). In this 925 situation, high tumor burden, high tumor grade, high Ki67 index, and uncontrolled symptoms 926 are major factors associated with worse prognosis {Assie, 2007 #114;Libe, 2015 #29}. 927 However, there is consensus that the kinetics of tumor growth might be also relevant, 928 particularly when making the decision for initiation of cytotoxic chemotherapy. However, this 929 parameter has not been formally assessed. Although a correlation of tumor growth and tumor 930 grade exists, it is not true for all tumors.
- 931 932

R.5.3. During follow-up, we recommend re-assessing prognosis at each evaluation, to guide treatment strategy (++OO).

935

936 <u>Reasoning</u>

After complete surgery, the major prognostic factor is whether there is any tumor recurrence.
At the time of recurrence the main prognostic factors are time between initial surgery and
recurrence, tumor burden and resectability {Datrice, 2012 #118;Erdogan, 2013 #55;Ettaieb,
2016 #117;Simon, 2017 #136}.

For patients with advanced disease, prognostic factors include Ki67 index, tumor burden,
general patient condition, and kinetics of tumor growth, as well as response to treatment.
Limited evidence is available, but these factors make clinical sense and are corroborated by
this panel's experience.

945 946

5.6. Methods and time interval for imaging and hormonal assessment duringfollow-up

949

950R.6.1. We recommend following patients with regular cross-sectional imaging of the
abdomen, pelvis and chest for disease recurrence or progression.

952 953 Reasoning

A majority of disease recurrence and progression occurs either loco-regionally, or with metastases to lung or liver and therefore should be identified by thoraco-abdomino-pelvic imaging. Bone metastases are infrequent and brain involvement is exceptional {Fassnacht, 2009 #56;Libe, 2015 #29;Burotto, 2015 #119}. In general, 18-FDG-PET/CT might provide additional information (see R.2.4.) particularly prior to any surgical intervention {Leboulleux, 2006 #120;Mackie, 2006 #121;Ardito, 2015 #122}. In addition, change in tracer uptake mightinform about disease evolution.

961 962

R.6.2. After complete resection, we suggest radiological imaging every 3 months for 2 years, then every 3-6 months for a further 3 years. The majority of the panel suggests continuation of follow-up imaging beyond 5 years, but surveillance should then be adapted.

967

968 <u>Reasoning</u>

969 There are no published studies that address specifically this issue. Therefore, the suggested 970 imaging interval is in accordance with the practice at many expert centers, and with 971 standards for other malignant tumors. In the experience of the panel few tumors with initial 972 curative surgery will recur after more than five years and therefore a 5-yr surveillance is likely 973 to include >90% of the ACC population that will experience disease recurrence. However, the 974 majority of the panel felt uncomfortable with the notion of complete cessation of imaging after 975 5 years and preferred for instance an annual imaging for another 5 years. After stopping 976 regular imaging, patients and primary care physicians should remain vigilant in terms of 977 potential symptoms or signs of late recurrences (see also R.6.4.).

978 979

983

R.6.3. For advanced ACC, we recommend surveillance based on prognostic factors, expected treatment efficacy and treatment-related toxicity, as well as the available alternative treatment options.

984 Reasoning

The imaging interval in advanced ACC depends on the ongoing treatment and the overall prognosis, but will usually be in 2-3 monthly intervals. For patients receiving mitotane alone, imaging intervals might be even more individualized (e.g. 2-5 months) based on tolerability and tumor kinetics. For patients undergoing loco-regional treatments, specific surveillance following procedures must be determined by the team performing these procedures, both to assess efficacy and adverse effects. For patients opting for entirely palliative management, without any anti-neoplastic therapy, no systematic imaging is advised.

992 993

994 R.6.4. In all patients, we recommend regular screening for hormone secretion.

995

996 <u>Reasoning</u>

997 Biochemical evaluation together with clinical evaluation fulfills two purposes: (i) it allows in a 998 few patients the early detection of recurrences and (ii) it also identifies patients that might 999 benefit from early anti-hormonal therapy. Biochemical evaluation should focus on steroid 1000 hormones or metabolites that were present at the time of diagnosis of the initial tumor. 1001 However, some panelists favored a more complete hormonal evaluation, because some 1002 tumors might change their steroid secretion pattern over time.

- 1003
- 1004

1005

1006 **5.7. Adjuvant therapy**

1007

1008**R.7.1.** For adrenal tumors with uncertain malignant potential, we recommend against1009adjuvant therapy (+OOO).

1010 <u>Reasoning:</u>

1011 In certain tumors it is difficult to define if the tumor is truly malignant (see R.4.4.). Since all 1012 adjuvant therapies are associated with potential toxicity, only patients with a definitive 1013 diagnosis of ACC should be considered for adjuvant treatment.

1014

R.7.2. We suggest adjuvant mitotane treatment in those patients without macroscopic residual tumor after surgery but who have a perceived high risk of recurrence (+OOO). However, we cannot suggest for or against adjuvant therapy for patients at low/moderate risk of recurrence (stage I-II, R0 resection and Ki67 ≤ 1019 10%) and adjuvant therapy options should be discussed on an individual basis.

1020 1021 Reasoning:

1022 The panel is in favor of offering mitotane to patients with high risk of recurrence (stage III, or 1023 R1 resection, or Ki67 >10%; see R.5.2.) despite the absence of completely convincing 1024 evidence (see section 4.3). The panel decided on the use of mitotane in the adjuvant setting 1025 based on three arguments: (i) the perceived effects {Terzolo, 2007 #33;Berruti, 2010 1026 #21;Berruti, 2017 #22;Bertherat, 2007 #82;Else, 2014 #125;Fassnacht, 2010 #57;Grubbs, 1027 2010 #191;Postlewait, 2016 #192} (acknowledging this is based on low guality evidence), 1028 see Figures 1A + B; (ii) published data showing a tumor response in ~20% of patients with 1029 advanced disease treated with mitotane {Baudin, 2001 #197;Else, 2014 #135;Hahner, 2005 1030 #64:Megerle, 2018 #294}; (iii) clinical experience of the panelists. For details on mitotane 1031 management see section 5.9.

Ki67 has emerged as the most powerful predictor of recurrence, and tumors with Ki67 ≤10%
might represent a subset of patients with a good prognosis. For these patients mitotane
might be considered overtreatment. For this subset of patients (<30% of all localized ACCs)
the ongoing ADIUVO trial, a prospective study where patients are randomized to adjuvant
mitotane vs. observation, will provide guidance in a few years.

1037 There is no clinical, histopathological, or molecular marker that reliably predicts response to 1038 mitotane although several markers have been proposed {Volante, 2012 #327;Ronchi, 2014 1039 #322}. A study showed that mitotane levels may influence patient outcome in adjuvant 1040 setting {Terzolo, 2013 #313} as it has been reported in advanced ACC. The secretory status 1041 of the tumor has a negative prognostic value but does not seem to influence response to 1042 treatment {Berruti, 2014 #35;Berruti, 2017 #22;Megerle, 2018 #294}.

- 1043 In patients who undergo surgery for recurrence of ACC but who have not previously had 1044 medical therapy, the decision on adjuvant mitotane should follow the same lines of 1045 reasoning.
- 1046 1047

1048R.7.3. Once the decision for mitotane treatment is established, we recommend1049starting mitotane as soon as clinically possible after surgery (+OOO).

1050 <u>Reasoning:</u>

1051 The ideal timing to start adjuvant mitotane is unknown; however, by analogy with other 1052 oncological adjuvant treatments we are convinced that starting mitotane within six weeks is 1053 ideal, and would not initiate the treatment later than 3 months. This reasoning is sound with 1054 the biological concept of adjuvant therapy in general, and with the latency of mitotane to reach effective levels and anti-tumor activity. However, no published data are available to
demonstrate the superiority of an early start of treatment or the lack of efficacy when started
later than 3 months.

1058 1059

R.7.4. In patients without recurrence who tolerate mitotane in an acceptable manner, we suggest to administer adjuvant mitotane for at least 2 years, but not longer than 5 years (+OOO).

1063 1064 Reasoning:

1065 The optimal duration of mitotane treatment is unknown and practice varies among different 1066 centers. Some members of the panel continue treatment for 3 to 5 years if tolerated {Terzolo, 1067 2014 #363}, while others discontinue after 2 to 3 years {Fassnacht, 2011 #61;Berruti, 2012 1068 #20;Else, 2014 #135}. Prognostic factors at diagnosis, patient compliance with treatment and 1069 plasma mitotane levels reached during treatment are factors that influence duration of 1070 treatment. Mitotane may possibly act as an oncostatic measure in those patients {Huang, 1071 2008 #350; Terzolo, 2009 #328}. However, the rate of recurrence 5 years after surgery is 1072 potentially too low to advise continuation of therapy treatment beyond this time point. 1073 Treatment-related toxicity, lack of experience in long-term administration are additional 1074 factors portending against indefinite treatment.

1075 1076

1077R.7.5. The panel did not come to a definitive consensus on adjuvant radiation therapy.1078However, we suggest against the routine use of radiation therapy in patients1079with stage I-II and R0 resection (+OOO). The panel suggests considering1080radiation in addition to mitotane therapy on an individualized basis therapy in1081patients with R1 or Rx resection or in stage III.

1082 1083 Reasoning:

1084 The systematic literature search indicated that radiation therapy is able to prevent local 1085 recurrence but does not significantly affect distant recurrences or overall survival {Else, 2014 1086 #125;Fassnacht, 2006 #126;Habra, 2012 #123;Polat, 2009 #73;Sabolch, 2015 #127;Sabolch, 2013 #124} (see section 4.3. and Figure 1). However, distant metastases 1087 1088 account for about 40-60% of tumor relapses {Berruti, 2017 #22;Amini, 2016 #157;Erdogan, 1089 2013 #55} and have large impact on the patient prognosis, and are more difficult to treat 1090 effectively. Conversely, prevention of the complications due to local recurrence argues in 1091 favor of radiation therapy. Adjuvant radiation therapy might be particularly reasonable in 1092 patients with R1 resection. This was already suggested by earlier studies, but also by a very 1093 recent study that was published after the meta-analysis associated with this report {Nelson, 1094 2018 #358}.

1095

1096 Radiation therapy is not advised for patients who experienced widespread tumor spillage 1097 during surgery. The combination of radiation therapy and mitotane is biologically sound 1098 {Cerquetti, 2008 #330;Cerquetti, 2010 #329} and possible but at the cost of greater toxicity 1099 (e.g. constitutional, gastrointestinal and liver toxicity). In addition, there is concern that 1100 radiation therapy may delay systemic therapy or prevent effective mitotane administration 1101 resulting in lower drug levels.

1103 1104 R.7.6. If adjuvant radiation therapy is administered, we recommend starting 1105 treatment as soon as clinically possible after surgery and to deliver radiation 1106 therapy at the dose of 50-60 Gy to the previous tumor bed in fractionated 1107 doses of approximately 2 Gy each (+OOO).

1108 <u>Reasoning:</u>

Radiation therapy was delivered following this scheme in previous observational studies
{Fassnacht, 2006 #126;Habra, 2012 #123;Sabolch, 2013 #124;Else, 2014 #125;Sabolch,
2015 #127} and lower dosage seems to be less effective {Polat, 2009 #73}.

1112 1113

1118

1114R.7.7. The panel did not come to a definitive consensus on adjuvant use of cytotoxic1115drugs. We suggest against the routine use of cytotoxic drugs in the adjuvant1116setting. However, the panel suggests considering adjuvant chemotherapy in1117selected patients with very high risk for recurrence.

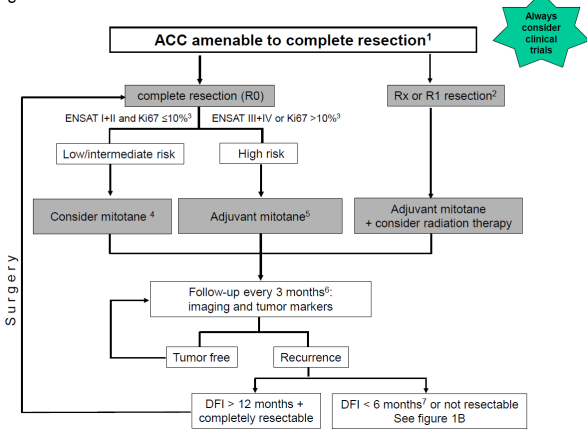
1119 <u>Reasoning:</u>

1120 Scant data are available on the use of cytotoxic drugs in an adjuvant setting and the studies 1121 did not control the results of treatment with a matched control group of untreated patients, or patients treated undergoing mitotane therapy {Khan, 2000 #296}. However, the majority of 1122 1123 panelists favors discussion of this option with patients with high risk of recurrence (ideally in 1124 the setting of clinical trials). Despite the lack of published data, some members of the panel 1125 are currently using cisplatin, with or without etoposide, in patients at perceived very high risk 1126 of recurrence (e.g. Ki67 >30%. large tumor thrombus in the vena cava, stage IV, or R1 1127 resection).

1128 In patients with R2 resection or tumor spillage, the same considerations for treatment of 1129 (locally) advanced disease should apply (see section 5.8.).

- 1130
- 1131

1132 Figure 3: Treatment for ACC amenable to complete resection



1134 1135

1136

- 1137 1138 1139 DFI disease-free interval between complete resection and recurrence
 - ¹ All patients with stage I+II and most patients with stage III should be amenable to radical resection. If complete resection is not feasible, consider neo-adjuvant treatment (e.g. cisplatin or EDP). In selected patients with stage IV and oligo-metastatic disease complete resection might be possible as well and should be aimed at.
 - ² In patients with R2 resection, consider re-surgery by an expert surgeon (see R.3.6) or see Figure 1B
- 1141 1142 ³ If Ki67 staining is not available, a low (<20 mitoses / 50 high power fields) or a high mitotic rate (> 20 mitoses / 1143 50 high power fields) may be used for risk stratification.
- 1144 ⁴ Individual decision (see R.7.2.). If possible enroll in clinical trial like ADIUVO (www.adiuvo-trial.org).
- 1145 ⁵ In some patients (e.g. Ki67 >30%. large tumor thrombus in the vena cava, stage IV, or R1 resection) consider additional cytotoxic therapy (e.g. 3-4 cycle of cisplatin + etoposide). 1146
- 1147 ⁶ After two years the time intervals are gradually extended.
- 1148 ⁷ If the disease-free interval is between 6 and 12 months or in patients with DFI > 12, in whom complete resection 1149 is not possible, an individual approach is required (see R.8.7.)

1150 **5.8. Treatment of recurrent and/or advanced ACC**

1151

1152 Clinical scenarios of patients with recurrent and/or advanced ACC are highly variable. 1153 Therefore, we try to provide recommendations for at least the most frequent presentations 1154 (see also Figure 4). Although a (small) proportion of patients experience a relatively long survival {Hermsen, 2008 #128;Fassnacht, 2009 #58;Libe, 2015 #29;Else, 2014 #135}, the 1155 1156 prognosis of advanced/metastatic ACC is generally limited. The goal of any therapy is to 1157 palliate symptoms and prolong survival. In this situation it is even more important than in 1158 other scenarios to tailor treatment on an individual basis taking into account the disease 1159 extent, the patient performance status and particularly the preference of the patient.

1160 1161

1162R.8.1. For patients presenting at time of initial diagnosis with limited intra-abdominal1163metastases we suggest surgical therapy if complete resection of all lesions1164seems feasible (+OOO). In case of limited extra-abdominal lesions, we suggest1165adrenal tumor resection in conjunction with therapy aiming at long-term tumor1166control of the other lesions (+OOO). In all patients, we recommend to start1167mitotane therapy as soon as clinically possible (+OOO).

1168

1169 <u>Reasoning:</u>

1170 Complete surgery is the best chance to reach long-term disease control although the 1171 likelihood of complete tumor removal in advanced ACC is low. If clinically possible, a single surgical approach should be planned. If a one-time surgical approach is impossible (e.g. due 1172 1173 to extra-abdominal metastases), other loco-regional approaches (see R.8.2) should be 1174 discussed within a multidisciplinary expert team and the patient on an individual basis. Local 1175 expertise and preference of the patient should be taken into account. Any initial treatment (surgery, local and/or medical therapy) should be initiated in a timely fashion (\leq 4-6 weeks 1176 1177 following initial diagnosis).

- 1178 In general, prognostic parameters (see R.5.2 + 5.3) should influence the overall treatment 1179 strategy. If the disease has an aggressive behavior (i.e. increase in tumor burden [e.g. 1180 increasing size of existing tumors or new metastasis] observed in subsequent imaging 1181 performed within a few weeks) systemic options (chemotherapy plus mitotane) may be 1182 favored. If partial responses or prolonged stabilization are then observed, surgery and/or 1183 additional loco-regional options might be particularly useful ("neo-adjuvant approach", see 1184 also R.8.3). This strategy could also be potentially advantageous in patients for whom tumor 1185 shrinkage might allow a more conservative surgical approach (i.e. patients in whom radical 1186 surgery would imply the complete or partial removal of neighboring organs such as kidney, 1187 spleen and part of the pancreas){Bednarski, 2014 #334}.
- 1188 These patients are at high risk for recurrence and therefore adjuvant mitotane seems to be 1189 justified {Wangberg, 2010 #361}. Addition of cytotoxic drugs might be a possible option 1190 (although data are lacking; see also R.7.7.).
- 1191 1192

1193**R.8.2.** The panel is convinced that in addition to surgery other local therapeutic1194measures (e.g. radiation therapy, radiofrequency ablation, cryoablation,1195microwave ablation, chemo-embolization) are of value for therapy of advanced1196ACC. We suggest individualization of the decision on the method of choice

1197based on the localization of the tumor lesion(s), local expertise, prognostic1198factors, and patient's preference (+OOO).

1199

1200 <u>Reasoning:</u>

Published data on local therapies in advanced ACC are very limited {Cazejust, 2010 #233;Ho, 2013 #234;Polat, 2009 #73;Wood, 2003 #235} and summarized in Appendix 6. However, the experience of many panelists provides additional support of efficacy of these local measures. Nevertheless, it is impossible to indicate which method is superior. Most important, the expertise of the local team in applying these methods should be taken into account when discussing this issue with patients in a shared decision-making process.

1207 1208

1209**R.8.3. We suggest against the routine use of adrenal surgery in case of widespread**1210metastatic disease at the time of first diagnosis (+OOO).

1211

1212 <u>Reasoning:</u> Despite the lack of large studies addressing this particular question, a majority of 1213 the panel agreed that patients with widespread and unresectable disease will usually not 1214 benefit from surgery. However, a few panelists suggested that adrenalectomy could be an 1215 option if technically possible.

- 1216 In patients who respond very well to systemic therapy, surgery should be considered at an 1217 appropriate time point; especially if complete resection becomes feasible ("neo-adjuvant 1218 approach"). However, the published evidence for such an approach is scant {Rangel, 2013 1219 #331;Bednarski, 2014 #334}.
- 1220 In selected cases (e.g. patients with severe hormone excess) debulking surgery might be an 1221 option, although anti-hormonal drugs (see R.10.1) should be considered here. In these 1222 cases, surgery might be especially reasonable if > 80% of the tumor burden can be removed 1223 safely. In patients with a poor clinical condition and significant localized metastatic burden, 1224 additional localized therapies (see R.8.2) may be considered as an alternative.
- 1225 1226

1230

1227R.8.4. In patients with advanced ACC at the time of diagnosis not qualifying for local1228treatment, we recommend either mitotane monotherapy or mitotane + EDP1229depending on prognostic parameters (+++O).

1231 <u>Reasoning:</u>

Mitotane is the treatment of choice for patients with advanced ACC (for details about the 1232 1233 management of mitotane see section 5.9). However, a very recent cohort study suggests that 1234 patients with metastastic disease at the time of primary diagnosis might not be the ideal 1235 candidates for mitotane monotherapy {Megerle, 2018 #294}. Furthermore, unfavorable 1236 prognostic parameters (e.g. high tumor burden, uncontrolled symptoms, high proliferative 1237 index, clinical evidence of a fast growing tumor) are important factors favoring a more 1238 aggressive/more rapidly active therapeutic approach. If more aggressive therapy is indicated, 1239 then the combination of EDP in addition to mitotane (EDP-M) is the most validated regimen 1240 {Fassnacht, 2012 #28}. EDP-M is the only treatment approach in ACC that is successfully 1241 evaluated in a randomized trial, the FIRM-ACT study. It has to be highlighted, however, that 1242 only progression-free survival was significantly improved in comparison to the alternative 1243 therapy (in this case streptozotocin plus mitotane; 5.0 vs. 2.1 months; HR 0.55; 95% CI 0.43

to 0.69; P<0.001)), whereas for overall survival the crossover design might have diluted the results (14.8 vs 12.0 months, HR 0.79; 95% CI, 0.61 to 1.02; P=0.07).

1246 The administration of EDP-M comes with risk of adverse events and it is important that the 1247 treatment will be administered by physicians with sufficient experience in oncology 1248 treatments. All cytotoxic drugs induce asthenia, nausea, vomiting and reversible 1249 myelotoxicity. In addition, etoposide might lead (among other adverse effects) to liver toxicity 1250 and reversible alopecia, doxorubicin to congestive heart failure and reversible alopecia; 1251 cisplatin to renal toxicity, otoxicity, peripheral neuropathy. In some patients, the risks might even outweigh the benefits (especially in patients with reduced performance status). If there 1252 1253 are concerns about the use of doxorubicin, cisplatin/carboplatin with or without etoposide (EP or P) might be an alternative option. Carboplatin may be an alternative to cisplatin, 1254 1255 particularly when cardiac or renal function is compromised. Again, in this cohort, loco-1256 regional treatment options may be particularly applicable.

Several studies have tried to find biomarkers that predict response to cytotoxic therapy in
ACC {Ronchi, 2009 #320;Malandrino, 2010 #335;Roca, 2017 #319;Laufs, 2018 #336}.
However, no reliable marker could be identified yet.

A few centers prefer the combination of etoposide and cisplatin (EP), because there is no single study proving that EDP is truly superior to EP. In patients with poor overall health cisplatin with mitotane may be an option. However, the evidence for etoposide + cisplatin or cisplatin alone is based only on small phase II studies {Bonacci, 1998 #199;Bukowski, 1993 #200;Williamson, 2000 #213}.

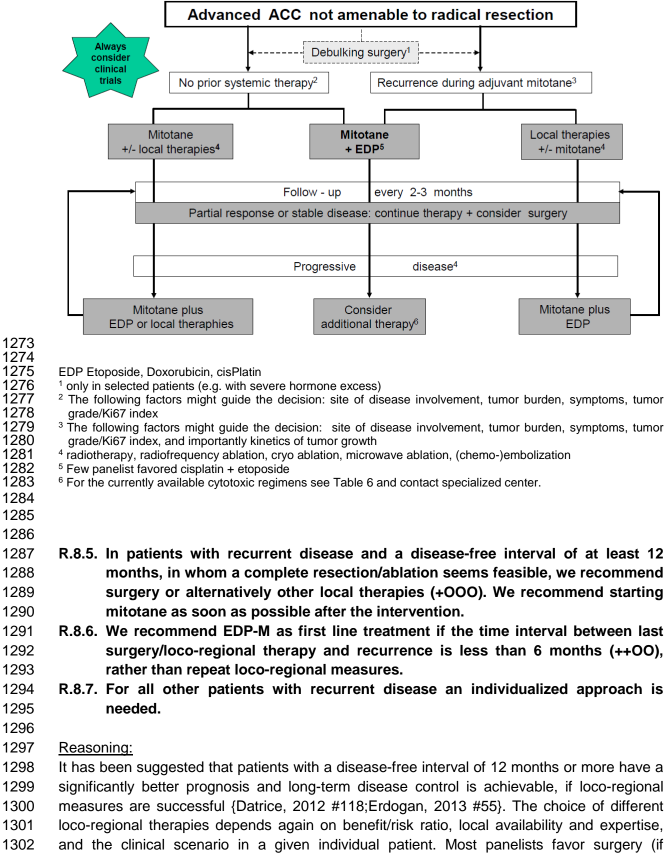
1265 There is limited evidence that standard chemotherapeutic agents may be more active in the 1266 presence of elevated mitotane concentrations {Bates, 1991 #295; Fassnacht, 2012 1267 #28;Sperone, 2010 #32}, but the panel is not in favor in delaying cytotoxic therapy for this 1268 reason for more than 14 days. Several centers start mitotane and cytotoxic therapy in 1269 parallel.

1271 Figure 4: Treatment of advanced ACC

1272

1270

Figure 4



1303 complete resection is feasible) followed by mitotane therapy.

1304 If the recurrence occurs during adjuvant mitotane therapy, additional measures could be 1305 considered. In patients with local recurrence, adjuvant radiation therapy after surgery should 1306 be discussed. In other scenarios, additional administration of cytotoxic drugs should be 1307 discussed with the patient, particularly when mitotane blood levels were in the recommended 1308 range > 14 mg/l.

Patients with early recurrence usually suffer from a very aggressive tumor, which most likely cannot be controlled by surgery or localized therapies. Decision-making should incorporate the concern that any local measure will only delay the administration of systemic therapy. Similar to the discussion to R.9.3, the FIRM-ACT data indicate EDP-M as the most effective form of therapy. An exception might be patients in whom incomplete initial surgery is the most likely cause for early progression. In these selected patients repeat surgery at an expert center might be an appropriate alternative (see R.3.6).

Patients with recurrence between 6 and 12 months after primary surgery usually have a poor prognosis and would, therefore, benefit from a more aggressive therapeutic approach (e.g. EDP-M). However, this decision should be discussed with the patient taking into account prognostic parameters (see section 5.5.), the feasibility of a R0 resection and patient's general condition. Patients with a disease-free interval > 12 months, in whom complete resection or loco-regional therapy is not feasible and who are currently not treated with mitotane, might be good candidates for mitotane monotherapy {Megerle, 2018 #294}.

1323 1324

1325**R.8.8.** In patients who progress under mitotane monotherapy, we recommend to add1326EDP (+++O).

1327

1328 Reasoning:

Mitotane is a slow-acting drug and in patients with rapidly progressing tumor, it might be too slow or not effective enough. In these patients, based on the FIRM-ACT data {Fassnacht, 2012 #28}, additional administration of EDP is the first choice (for alternatives see Reasoning R.8.4.). However, if the tumor burden is limited despite obvious progression, another 2-3 months mitotane monotherapy could also be justified, particularly if adequate mitotane levels have not been achieved. In these cases, additional loco-regional options should be considered.

1336 1337

1343

1338R.8.9. In patients who respond to medical therapy (including achievement of long-1339term stable disease), we suggest re-considering local measures aiming at long-1340term tumor control. Such an approach could be also considered in patients1341attaining a generally good control of the disease, in which a limited number of1342lesions are progressing.

1344 <u>Reasoning:</u>

1345 In some patients, in whom long-term disease control could be achieved, loco-regional 1346 measures (in addition to ongoing medical therapy) might be able to reach complete 1347 remission or at least significantly reduce tumor burden {Berruti, 2005 #24}. In patients with 1348 "mixed responses"; e.g. progressive disease limited to few lesions, loco-regional options 1349 might be reasonable to add to the ongoing medical therapy.

1350

1351

1352**R.8.10.** In patients who progress under EDP-M we suggest considering additional1353therapies including clinical trials on an individual basis (+OOO).

- 1354
- 1355 <u>Reasoning:</u>

1356 Several drugs and drug combinations have been tested in advanced ACC. However, except 1357 EDP-M none of them has been successfully evaluated in large randomized trials. Figure 2 outlines the outcomes of the different approaches. However, this figure has to be interpreted 1358 1359 with great caution, because differences in the characteristics of the patients included in the different cohorts preclude direct comparison between studies. Therefore, it is not possible to 1360 1361 draw definitive conclusions. Due to the limited treatment options, the panel clearly favors 1362 enrollment of patients with progressing tumors in clinical trials investigating experimental 1363 therapies including phase I trials. However, the panel felt that despite the lack of convincing 1364 data, some guidance might be helpful for patients that cannot be enrolled in clinical trials 1365 (Table 6). Beyond cisplatin-based therapies, the two reasonably well-studied second-line cytotoxic regimens are gemcitabine + capecitabine (+/- mitotane) {Henning, 2017 1366 1367 #215;Sperone, 2010 #32} and streptozotocin + mitotane {Khan, 2000 #296;Fassnacht, 2012 1368 #28}. However, objective response rates are clearly below 10% and median progression-free 1369 survival (PFS) is generally <4 months, but a few patients with long-term disease control and even complete responses in single patients are described. Nevertheless, a few panelists 1370 1371 argued against the use of streptozotocin, because median PFS in the FIRM-ACT trial was 1372 only two months {Fassnacht, 2012 #28}. As for EDP, these cytotoxic drugs should be 1373 administered only by physicians experienced with chemotherapy. Typical adverse effects of 1374 streptozotocin are nausea, vomiting, diarrhea, renal and liver toxicity and of the association 1375 gemcitabine and capecitabine nausea, vomiting and reversible myelotoxicity.

Loco-regional measures can be particularly useful when progression is limited, or only
affects limited areas (e.g. single organs). In these cases, such localized therapies (see
R.8.2) might be able to provide higher response rates for these specific organ/tissue areas
than second line systemic options.

- 1380 Several tyrosine kinase inhibitors have been investigated in advanced ACC {Berruti, 2012 1381 #23:Fassnacht, 2015 #27:Kroiss, 2012 #207:O'Sullivan, 2014 #209}, but the results were 1382 largely disappointing. However, in retrospect, drug efficacy could have been hampered by 1383 increased metabolism of the TKI due to mitotane-induced CYP3A4 activity. Nevertheless, 1384 currently no specific TKI can be suggested for the treatment of advanced ACC. Targeting the 1385 IGF2/IGF receptor signaling pathway was pathophysiologically a very promising approach and initial small studies suggested some efficacy {Almeida, 2008 #303;Boulle, 1998 1386 #304;Gicquel, 1994 #305;Giordano, 2003 #306;Weber, 2000 #307;Haluska, 2010 1387 #204; Jones, 2015 #299; Lerario, 2014 #216; Naing, 2011 #301; Naing, 2013 #208}. However, 1388 1389 the large placebo-controlled phase III GALACCTIC trial demonstrated that the IGF1R 1390 inhibitor linsitinib did not improve progression-free or overall survival {Fassnacht, 2015 #27}. 1391 Therefore, monotherapy with drugs targeting this pathway are not reasonable for therapy in an unselected patient population. 1392
- 1393
- 1394 1395

Table 6: Systemic therapies for recurrent / advanced ACC

First-line therapies (see text for details)

• Surgery +/- other local measures (see R.8.1 and R.8.4)

•	Mitotane monotherapy - details on the management see section 5.9.
•	Etoposide, Doxorubicin and Cisplatin (EDP) plus Mitotane (EDP/M) {Fassnacht, 2012 #28}
	every 28 days:
	day 1 40mg/m ² doxorubicin (D)
	day 2 100mg/m ² etoposide (E)
	day 3+4 100mg/m ² etoposide (E) + 40mg/m ² cisplatin (P)
	plus oral mitotane aiming at a blood level between 14-20mg/l.
	In patients unfit for the EDP-M regimen, (E)P-M may constitute a reasonable alternative. Every 28 days
	day 1 100mg/m ² etoposide (E)
	day 2+3 100mg/m ² etoposide (E) + 40mg/m ² cisplatin (P)
<u>Ad</u>	litional therapeutic options
	litional therapeutic options Consider enrollment of patients in clinical trials (www.clinicaltrial.gov)
<u>Ad</u> •	
	Consider enrollment of patients in clinical trials (www.clinicaltrial.gov) Consider loco-regional therapies
	Consider enrollment of patients in clinical trials (www.clinicaltrial.gov)
	Consider enrollment of patients in clinical trials (www.clinicaltrial.gov) Consider loco-regional therapies Gemcitabine plus capecitabine {Henning, 2017 #215;Sperone, 2010 #32}
	Consider enrollment of patients in clinical trials (www.clinicaltrial.gov) Consider loco-regional therapies Gemcitabine plus capecitabine {Henning, 2017 #215;Sperone, 2010 #32} 800 mg/m ² gemcitabine on day 1 and 8 (repeated every 3 weeks)
	Consider enrollment of patients in clinical trials (www.clinicaltrial.gov) Consider loco-regional therapies Gemcitabine plus capecitabine {Henning, 2017 #215;Sperone, 2010 #32} 800 mg/m ² gemcitabine on day 1 and 8 (repeated every 3 weeks) 1,500 mg capecitabine orally per day in a continuous fashion Mitotane can be continued (individualized decision) Streptozotocin plus Mitotane (Sz/M) {Fassnacht, 2012 #28}
	Consider enrollment of patients in clinical trials (www.clinicaltrial.gov) Consider loco-regional therapies Gemcitabine plus capecitabine {Henning, 2017 #215;Sperone, 2010 #32} 800 mg/m ² gemcitabine on day 1 and 8 (repeated every 3 weeks) 1,500 mg capecitabine orally per day in a continuous fashion Mitotane can be continued (individualized decision) Streptozotocin plus Mitotane (Sz/M) {Fassnacht, 2012 #28} induction: day 1-5: 1g Sz/d
	Consider enrollment of patients in clinical trials (www.clinicaltrial.gov) Consider loco-regional therapies Gemcitabine plus capecitabine {Henning, 2017 #215;Sperone, 2010 #32} 800 mg/m ² gemcitabine on day 1 and 8 (repeated every 3 weeks) 1,500 mg capecitabine orally per day in a continuous fashion Mitotane can be continued (individualized decision) Streptozotocin plus Mitotane (Sz/M) {Fassnacht, 2012 #28}

1396

1397

1398**R.8.11.** The optimal timing of mitotane discontinuation is currently unknown and the1399panel could not come to a specific recommendation on this issue.

1400

1401 <u>Reasoning:</u>

1402 A recent cohort study reported that discontinuation of mitotane should be considered in 1403 patients who experienced progressive disease after one year of mitotane therapy {Vezzosi, 1404 2018 #308}. Part of the panel considers mitotane discontinuation when there is progressive 1405 disease despite mitotane blood levels above 14 mg/L while others often continue mitotane 1406 indefinitely in their practice. Tolerability of treatment is an important issue to consider in this 1407 decision. Moreover, it has to be considered that CYP3A4 induction by mitotane can greatly 1408 enhance metabolism of many drugs {Kroiss, 2011 #72}, including a number of experimental 1409 anti-ACC compounds, and so potentially limit their effectiveness.

1410 1411

5.9. Special considerations on mitotane

- 1412 If mitotane therapy is started (independent of the clinical scenario) the following issues have 1413 to be considered.
- 1414

1415R.9.1. We recommend starting therapy with mitotane in an escalating regimen1416depending on the performance status of the patient as well as the tolerability in1417the first weeks.

1418

1419 <u>Reasoning</u>

1420 There are different regimens to administer mitotane, but none of them has been proven to be 1421 superior. In patients with good performance status some panelists use a high starting dose 1422 approach: mitotane is administered at a starting dose of 1.5 g/day and if well-tolerated from a 1423 gastrointestinal perspective the dose is increased on day two to 3 g/day, on day three to 4.5 1424 g/day, and on day four to 6 g/day {Faggiano, 2006 #309;Mauclere-Denost, 2011 #310}. This 1425 dosage will be administered until first mitotane blood level is assessed. In this high dose 1426 regimen, it is strongly recommended to measure mitotane blood levels 2-3 weeks after 1427 initiation of therapy. Afterwards dosage will be adjusted according to blood concentrations 1428 and tolerability. Other panelists prefer a low starting dose approach. With this approach, 1429 mitotane is administered at a starting dose of 1 g/day and increased when there is good 1430 gastrointestinal tolerance every 3 days by 0.5 g up to a total dose of 3.0 - 4.0 g/day and then 1431 adjusted according to blood concentrations and tolerability {Terzolo, 2000 #311;Terzolo, 1432 2008 #364;Terzolo, 2014 #363}.

- 1433 In a formal comparative pharmacokinetic study, the high-dose starting regimen led to slightly 1434 higher mitotane plasma levels within 12 weeks of treatment, and more patients reached the 1435 target level of 14 mg/L {Kerkhofs, 2013 #38}. However, these results were not statistically 1436 significant due to lack of power. Beyond these two regimens, there is a variety of other 1437 possibilities and choice depends on personal practice, clinical scenario and patient 1438 conditions.
- 1439 Mitotane is a lipophilic drug and is supposed to be better absorbed from the gut with a high 1440 fat content of the diet, e.g. with milk or chocolate. {Moolenaar, 1981 #374}. In case of limited 1441 gastrointestinal tolerance, symptomatic treatments of nausea and or diarrhea may be 1442 proposed.
- 1443

1444R.9.2. We recommend monitoring of blood concentration of mitotane. The general aim1445is to reach a mitotane blood level above 14 mg/L (+OOO).

1446

1447 <u>Reasoning</u>

As long as mitotane plasma levels are increasing and have not yet reached a plateau at >1449 >14mg/L, mitotane plasma levels will be assessed every 3-4 weeks. Mitotane plasma level determination is best done as morning trough sampling, at least 12 hours after the last dose, preventing false high levels {Kerkhofs, 2014 #351}. When mitotane plasma levels have reached a plateau, it is usually sufficient to measure blood levels every 6-12 weeks.

1453 Usually it takes several weeks (sometimes months) to reach mitotane levels > 14 mg/L. As 1454 long as the concentration is < 14 mg/L it is reasonable to continue to increase the dosage if 1455 this is tolerated by the patient. Due to slow pharmacokinetic characteristics, the dose of 1456 mitotane can be reduced in most patients as soon as a plasma level of > 14mg/L is reached. 1457 Over time, mitotane dosage will be titrated to the best tolerable dose while maintaining a 1458 plasma level >14mg/L. Most patients experience adverse effects to a certain extent and 1459 these usually correlate with the plasma mitotane level (although there is major inter-individual 1460 variability) (see Table 7). However, some gastrointestinal adverse effects (like diarrhea) 1461 seem to correlate more with the oral dosage than with the plasma level and occur more 1462 frequently in the first phase of treatment {Terzolo, 2000 #311;Allolio, 2006 #236;Daffara, 1463 2008 #25;Terzolo, 2008 #364;Terzolo, 2014 #363}. Several studies {van Slooten, 1984 1464 #312;Baudin, 2001 #197;Haak, 1994 #202} have shown that CNS-related adverse events in particular occur more frequently when the plasma mitotane is > 20 mg/L. Therefore, many 1465 1466 experts recommend aiming to keep plasma concentrations below 20 mg/L. However, it can 1467 be speculated that higher plasma levels may also be associated with better clinical efficacy. 1468 Furthermore, some patients do not experience relevant adverse events even at plasma 1469 levels well above 20 mg/L. Regarding the lower limit it has to be acknowledged that in at 1470 least a few patients objective responses are seen even though plasma levels of >14 mg/l 1471 were not achieved {Megerle, 2018 #294}. Therefore, some panelists favored a target range 1472 of plasma mitotane of 8-30 mg/L, whereas others aim at an individualized target level of 1473 mitotane.

1475 Most studies addressing plasma mitotane levels analyze patients with advanced disease. 1476 However, there is one study suggesting that the same target level is also reasonable for the 1477 adjuvant setting {Terzolo, 2013 #313}. Therefore, the panel is in favor to use the same 1478 approach for both patient groups.

1479 1480

1474

1481R.9.3. We recommend glucocorticoid replacement in all patients treated with mitotane1482(except those with ongoing cortisol excess). We suggest to using1483hydrocortisone/cortisone acetate for this purpose. Due to increased steroid1484clearance and increase cortisol-binding globulin at least twice the standard1485replacement dose is usually required.

1486

1487 <u>Reasoning</u>

1488 A possible strategy is to start concomitant treatment on day one of mitotane treatment with 1489 hydrocortisone 20 mg/d. Alternatively, patients can be instructed to start hydrocortisone later 1490 (e.g. after 2-3 weeks or in case they experience adrenal insufficiency), because impairment 1491 of glucocorticoid effectiveness is rarely observed within the first few weeks. Due to the 1492 increased clearance and increased cortisol-binding globulin {Daffara, 2008 #25;Chortis, 2013 1493 #52:Reimondo, 2017 #349:Kerkhofs, 2015 #39} with increasing mitotane plasma levels and 1494 based on clinical symptoms, the total hydrocortisone replacement dose will usually increased 1495 to a typical total daily dose of 50 mg in two or three divided doses. However, some patients 1496 require daily dosages up to 100 mg. There is no reliable laboratory marker to guide the optimal dosage of hydrocortisone {Reimondo, 2017 #349}, which has to be based on clinical 1497 1498 judgment similar to the management of patients with adrenal insufficiency (Bornstein, 2016) #314}. Mitotane-induced increase in cortisol-binding globulin may confound interpretation of 1499 1500 serum cortisol measurement. The measurement of free cortisol may offer additional 1501 information, but more studies are required to clarify the value of this method {Alexandraki, 1502 2010 #325}. Some panelists measure plasma ACTH and use ACTH levels more than 2-fold 1503 of the upper limit of normal as evidence for insufficient glucocorticoid replacement. Other 1504 centers prefer a combined measurement of plasma ACTH and 24-hour urine free cortisol 1505 levels to assess adequacy of and optimize glucocorticoid replacement for patients receiving mitotane. However, when urinary cortisol is measured by immunoassays, interference bycortisol metabolites induced by mitotane might occur.

1508 In case of acute adverse events and/or hospital admission, patients should be treated 1509 intravenously with high-dose hydrocortisone (e.g. 100 mg TID) until resolution of symptoms.

Some patients experience symptoms and signs of insufficient mineralocorticoid activity (hyperkalemia, hyponatremia, hypotension, decreased wellbeing) despite full-dose substitution with hydrocortisone. In these patients, addition of fludrocortisone should be considered. Clinical judgment, electrolytes, and plasma renin concentration can be used for decision making whether to start fludrocortisone {Allolio, 2006 #236;Daffara, 2008 #25;Terzolo, 2008 #364;Terzolo, 2014 #363}.

1516 1517

1521

R.9.4. We recommend regular monitoring of mitotane-induced adverse effects (Table 7) and to treat them appropriately (Table 8). To increase tolerability of mitotane, we suggest starting supportive therapy ideally before severe toxicity occurs.

- 1522 Reasoning
- 1523 In addition to adrenal insufficiency (see R.9.3.) mitotane treatment comes with a plethora of 1524 potential adverse events {Daffara, 2008 #25}(Table 2). Therefore, it is important to evaluate 1525 the patients regularly (e.g. in the first 6 months every 3-4 weeks, thereafter every 6-12 1526 weeks).
- 1527 Gastrointestinal adverse effects are frequent, particularly in the first months of therapy. 1528 Supportive therapy should include antiemetic and anti-diarrheal medication, as needed. 1529 Some centers even start supportive therapy at initiation of mitotane therapy. However, one 1530 has to be aware that nausea may also be a sign of adrenal insufficiency that needs 1531 recognition and appropriate treatment. Nevertheless, it should be emphasized that despite 1532 optimization of dosing schedules, the key factor influencing build-up of appropriate mitotane 1533 plasma levels is patient tolerability, so efforts should be made in order to optimize this.
- 1534 In case of central nervous system (CNS) adverse effects grade 2 (moderate) and/or gastro-1535 intestinal adverse effects grade 3 (severe, but not life-threatening), mitotane dose should be 1536 reduced by 1-1.5 gram/day. In case of CNS severe, but not life-threatening (grade 3) adverse 1537 effects or any relevant grade 4 toxicity (life-threatening), and/or increase of liver enzymes >5 1538 times baseline (except GGT), mitotane should be interrupted until significant improvement of 1539 symptoms occurs and be restarted at 50–75% of the last dose.
- Assessment of thyroid hormone status (TSH, FT4, every 3 months) is advised as mitotane may induce a clinical picture similar to central hypothyroidism {Daffara, 2008 #25;Russo, 2016 #365}, possibly through a direct effect on the pituitary gland or induction of thyroid hormone metabolism. Replacement therapy with levothyroxine can be considered for these patients.
- 1545 In men with signs of hypogonadism, assessment of testosterone and sex hormone-binding 1546 globulin levels is warranted, as hypogonadism is common {Daffara, 2008 #25}. Mitotane-1547 induced increase in SHBG may confound interpretation of testosterone measurement. 1548 Testosterone supplementation may be considered in patients with low testosterone and 1549 symptoms of hypogonadism, but inhibition of 5-α reductase might prevent full activity of 1550 testosterone {Chortis, 2013 #52}.
- 1551 Ovarian steroid synthesis is less affected but women in childbearing age treated with 1552 mitotane may develop multiple, and sometimes huge, ovarian cysts that may be painful and 1553 sometimes require treatment.

1554 Cholesterol levels very frequently increase during mitotane treatment {Tada, 2014 #332}. 1555 Hypercholesterolemia can be treated with statin therapy using agents not metabolized by CYP3A4 (e.g. rosuvastatine or pravastatine). However, HDL cholesterol is usually also 1556 1557 elevated significantly and this should be taken in consideration. Thus, statin therapy might only be beneficial in selected patients (e.g. with good prognosis in an adjuvant setting, high 1558 1559 LDL cholesterol and additional high cardiovascular risk factors). Therefore, an indivdual 1560 decision making regarding the benefits of any lipid lowering therapy is necessary.

Psychological and social aspects of treatment should not be neglected, i.e., professional 1561 1562 counseling may be warranted. Follow-up on patient's well-being may be performed by 1563 guestionnaire-based assessment of toxicity upon the start of the treatment and by repeating 1564 this assessment every 3 months.

1565 1566

R.9.5. We recommend being aware of significant drug interactions of mitotane (e.g. 1567 due to strong induction of CYP3A4). All concomitant medication should be 1568 checked for CYP3A4 interactions and substituted for an alternative if necessary 1569 1570 and available. Other care-providers should be advised not to initiate other drug 1571 therapies without consultation.

1573 Reasoning

1574 A comprehensive (but not exhaustive) summary of relevant drug interactions with mitotane is 1575 provided in Kroiss et al. {Kroiss, 2011 #72} and in the Appendix 7.

1576 1577

1572

1578

1579

1580

1581

Table 7: Adverse effects during mitotane treatment*

Adverse Effect	
Gastrointestinal: nausea, vomiting, diarrhea, anorexia	very common
Adrenal insufficiency	very common
CNS: lethargy, somnolence, vertigo, ataxia	common
Confusion, depression, dizziness, decreased memory	common
Increase of hepatic enzymes (in particular gamma-GT)	very common
Liver failure	rare
Hepatic microsomal enzyme induction with	
increased metabolism of glucocorticoids and other steroids	very common
and barbiturates, phenytoin, warfarin, and many other drugs (see Appendix 7)	common
Increase in hormone-binding globulins (CBG, SHBG, TBG, etc.)	very common
Disturbance of thyroid parameters (interference with binding of T4 to TBG, total	very common
T4↓, free T4↓, TSH↓)	
Hypercholesterolemia, hypertriglyceridemia	very common
Gynecomastia	very common
Skin rash	common
Primary hypogonadism in men	common
Prolonged bleeding time	common
Leucopenia	common
Thrombocytopenia, anemia	rare
Autoimmune hepatitis	rare
Cardiovascular: hypertension	not known
Ocular: blurred or double vision, toxic retinopathy, cataract, macular edema	not known
Hemorrhagic cystitis	not known

- *modified by the authors based on information published by the European Medicine Agency 1582 (EMA):
- 1583 http://www.ema.europa.eu/ema/index.jsp?curl=pages/medicines/human/medicines/000521/human 1584 _med_000895.jsp&mid=WC0b01ac058001d124 and clinical experience

1585Frequency is defined according to the following convention: very common ($\geq 1/10$), common1586($\geq 1/100$ to <1/10), uncommon ($\geq 1/1,000$ to <1/100), rare ($\geq 1/10,000$ to <1/1,000), very rare</td>1587(<1/10,000), Not known (cannot be estimated from the available data)</td>

Table 8: Monitoring during mitotane treatment

Parameter	Interval	Comment
Recommended mo	onitoring	
Mitotane blood level	Every 3-4 weeks, as soon as plateau of blood level is reached every 2-3 months	Target blood level > 14 mg/L (details see R.9.2)
GOT, GPT,	Initially every 3-4 weeks,	GGT is invariably elevated without clinical consequences. If othe
bilirubin, (gGT)	after 6 months every 2-3 months	liver enzymes are rapidly increasing (> 5-fold of baseline), there risk of liver failure: interrupt mitotane
Blood count	Initially after 3-4 weeks, then every 3-4 months	Check for rare and in most cases not significant leucopenia, thrombocytopenia, and anemia
Suggested monito	oring	
ACTH	Suspected glucocorticoid deficiency or excess	Glucocorticoid status is difficult to determine Target: ACTH in the normal range or slightly above
TSH, fT4	Every 3 – 4 months	Disturbance of thyroid hormones is frequent. Thyroid hormone replacement is only recommended in patients with clinical symptoms of hypothyroidism
Renin	Every 6 months	If renin \uparrow and clinical symptoms of hypoaldosteronism are preser add fludrocortisone
Cholesterol (HDL, LDL)	Every 3-4 months (in adjuvant setting)	If LDL / HDL cholesterol $\uparrow\uparrow$ consider treatment with statins in selected cases.
Testosterone and SHBG in men	Every 3-4 months (in adjuvant setting)	If testosterone is low and clinical symptoms of hypogonadism are present add testosterone

1596 **5.10. Other supportive therapies**

1597

1600

1593 1594 1595

1591 1592

1598**R.10.1.** We recommend medical therapy to control hormone excess in all patients with1599clinically relevant hormone-producing ACC.

1601 Reasoning

1602 Overt glucocorticoid excess causes significant morbidity, such as diabetes, osteoporosis, 1603 muscle weakness and immunosuppression, conditions that can impact quality of life and 1604 increase mortality. Mitotane is effective in controlling adrenocortical hormone excess 1605 syndromes, but its efficacy is delayed by several weeks. In general, mild hormone secretion can be effectively managed by mitotane alone. However, severe Cushing syndrome needs a 1606 more rapid control. Furthermore, these patients should receive appropriate anticoagulation 1607 and also pneumocystis directed antibiotic prophylaxis until cortisol levels are safely controlled 1608 1609 {Nieman, 2015 #133}. In selected patients, surgery might even be postponed for few weeks 1610 until Cushing's syndrome is partly under control with the use of rapid agents inhibiting 1611 steroidogenesis (i.e. metyrapone). However, some panelists argued that surgery might be 1612 the fastest way to control severe hypercortisolism.

- Available steroidogenic enzyme inhibitors and steroid receptor antagonists are able to attain quick reduction of cortisol effects. Anti-hormonal agents can be initiated together with mitotane. Once therapeutic mitotane levels are established, anti-steroidogenic action is also maximized, and other anti-hormonal drugs can be reduced guided by tolerability, symptoms and biochemical measurements. If possible doses should be titrated to normalization of hormone levels, or in the case of receptor antagonists to improved well-being, accepting that assessment of this can be challenging in cancer patients.
- 1620 Despite the lack of comparative studies, the majority of panel members considers that 1621 metyrapone is the first therapeutic choice for the management advanced ACC patients with
- 1622 severe Cushing syndrome. The drug is well tolerated and can be safely administered in

1623 association with mitotane and cytotoxic chemotherapy {Claps, 2017 #131}. Moreover, its 1624 metabolism and elimination are not altered by concomitant mitotane. Ketoconazole an 1625 inhibitor of several key cytochrome P450 (CYP) enzymes involved in multiple steps of 1626 steroidogenesis in the adrenal cortex, is another alternative, but often less effective than 1627 metyrapone and requires regular monitoring of liver function tests. Its advantage is that it 1628 also inhibits androgen production. Ketoconazole should be avoided at initiation of mitotane 1629 therapy because both substances are potentially hepatotoxic and it will be difficult to attribute 1630 the hepatotoxicity to one or the other drug. Hypercortisolemia can also be treated with 1631 mifepristone, a glucocorticoid antagonist, but dosing is based on clinical judgement as 1632 cortisol levels remain elevated or rise further on therapy {Castinetti, 2009 #132}. Moreover, 1633 the high circulating cortisol levels when on mifepristone may cause mineralocorticoid effects, 1634 including hypertension and hypokalemia that necessitate treatment with high doses of 1635 spironolactone. Patients treated with enzyme inhibitors or receptor antagonists need to be 1636 educated about symptoms and signs of adrenal insufficiency. All patients at risk for adrenal 1637 insufficiency need to be supplied with emergency medication and instructions. Intravenous 1638 etomidate can be used for seriously ill patients with severe hypercortisolemia who cannot 1639 take oral medication.

1640 In the management of severe Cushing's syndrome, locoregional options (see R.8.2.) should 1641 also be discussed, in selected cases.

- 1642 Androgen excess in women can impact quality of life due to hirsutism and virilization. It can 1643 be treated with androgen receptor antagonists, such as bicalutamide, flutamide, or 1644 spironolactone.
- 1645 Only a small fraction of all tumors produce aldosterone, leading to hypertension and 1646 hypokalemia. Mineralocorticoid excess is best treated with mineralocorticoid receptor 1647 antagonists, such as spironolactone or eplerenone. However, patients with severe Cushing's 1648 syndrome may also experience hypokalemia, related to mineralocorticoid receptor activation. 1649 In case of severe hypokalemia, spironolactone and epithelial sodium channel inhibitors such 1650 as amiloride can be used, potentially at high doses, along with potassium supplementation. 1651 In such cases, frequent serum electrolyte measurement, initially several times a week, are 1652 mandatory, as there is a risk of rapid occurrences of hyperkalemia and hyponatremia.
- 1653 In the rare situation of estradiol production by tumors in male patients, therapy with estrogen 1654 receptor antagonists or aromatase inhibitors could be considered.
- 1655 1656

1657**R.10.2.** We recommend therapy with anti-resorptive treatment in patients with bone1658metastasis.

1659

1660 <u>Reasoning</u>

1661 Bone metastasis in cancer patients are associated with poor quality of life due to bone pain 1662 and increased risk of adverse skeletal-related events (SREs) such as pathological fractures, 1663 spinal cord compression and hypercalcemia. Several randomized phase III trials have 1664 demonstrated that bone resorption inhibitors such as bisphosphonates and denosumab are 1665 efficacious in the prevention of skeletal-related events in patients with bone metastasis from 1666 breast, prostate, lung and others primary malignancies. No data are available for ACC 1667 patients. However, based on these results, it has become general practice to treat patients with any kind of bone metastasis with anti-resorptive therapies. The administration of 1668 1669 denosumab or bisphosphonates in 'oncological doses' in association with calcium intake and 1670 vitamin D supplementation are therefore advisable in ACC patients with metastatic bone

disease, with the aim to prevent adverse skeletal-related events and improve control of bonepain.

1673 In patients with ACC with Cushing's syndrome that cannot be otherwise controlled anti-1674 resorptive treatment, using 'anti-osteoporotic doses', should be considered, because it is well 1675 established that glucocorticoid-excess increases the risk of osteoporotic fractures. Since 1676 fracture risk declines rapidly after lowering excess cortisol, or antagonizing its effects, anti-1677 osteoporotic therapies are usually not required once cortisol secretion is controlled (either by 1678 surgery or medical therapy).

1679

1680

1683

1681R.10.3. We recommend palliative radiation for symptom palliation in1682advanced/metastatic ACC patients

1684 Reasoning

1685 Palliative radiation therapy is a commonly utilized intervention for symptom relief among 1686 patients with metastatic cancer. Two schedules of irradiation are commonly used, which 1687 include 8 Gy in a single fraction or 30 Gy in ten fractions. This treatment modality is highly effective in achieving relief of symptoms arising from bone metastases, with positive 1688 responses in up to 50% - 90% of cancer patients {Chow, 2012 #355;Pin, 2018 #356}. Painful 1689 bone metastases are, therefore, the main indication of palliative radiation in metastatic ACC 1690 1691 patients {Polat, 2009 #73}. Other indications are symptomatic recurrences, severe mass 1692 effect and the rare case of brain metastases.

1693 1694

1697

1695R.10.4. We recommend integrating palliative care into standard oncology care for all1696patients with advanced ACC

1698 <u>Reasoning</u>

According to the WHO palliative care is defined as 'an approach that improves the quality of life of patients and their families facing the problems associated with life-threatening illness, through the prevention and relief of suffering by means of early identification and impeccable assessment and treatment of pain and other problems, physical, psychosocial and spiritual' (WHO: WHO definition of palliative care. http://www.who.int/cancer/palliative/definition/en/).

1704 As previously stated, the goal of care for metastasized ACC is to obtain long-term disease 1705 control and prolong patient survival. Although prognosis of patients with advanced ACC is often poor, there is a patient subset destined to obtain a relatively long survival, while treated 1706 1707 with antineoplastic therapies. The needs of patients with cancer and their families have 1708 changed over time. According to the ASCO guidelines the best model to manage metastatic 1709 patients is to integrate palliative care early in the course of the disease and throughout the 1710 trajectory of care, extending to long-term survivorship as well as end-of-life (hospice) care. In 1711 this integrated approach the primary endocrinologists and oncologists focus on the primary 1712 oncologic disease, and the palliative care team addresses the majority of the patient's 1713 physical and psychological concerns. The team plans all therapy aiming to integrate patient 1714 wishes and employ treatment options balancing quality of life and increased survival with 1715 therapy associated risks and complications {Ferrell, 2017 #316}.

- 1716
- 1717
- 1718

R.10.5. We suggest counseling for fertility protection in female patients in reproductive age. Fertility counseling should not only be restricted to patients undergoing cytotoxic chemotherapy, but also given to patients who plan to embark on mitotane therapy.

1723 1724 Reasoning

A considerable proportion of patients are diagnosed with ACC during their reproductive years. Several drugs used to treat ACC harbor significant risk for impairment of fertility or the exact risks are unknown (e.g. mitotane). On the other hand, in recent years several treatment options for preservation of fertility have been introduced. However, none of them has gained general acceptance. Therefore, we just advise to discuss this topic with every patient. This discussion should include the consideration given in section 5.12. on pregnancy and ACC in general.

1732

1733 1734

1735 5.11. Genetic counseling

1736

1740

1737R.11.1. For adults with ACC, we recommend at least a basic clinical genetic1738evaluation, exploring personal and family history for evidence of a hereditary1739predisposition syndrome.

1741 <u>Reasoning</u>

1742 The detection of germline mutations impacts on the clinical care and surveillance of index 1743 patients and offers the possibility to identify at risk family members. Probably, up to 5% of 1744 adult ACC arise in patients with germline TP53 mutations {Herrmann, 2012 #287;Raymond, 1745 2013 #289;Waldmann, 2012 #77} and about 3% of all ACC patients have an underlying diagnosis of Lynch syndrome {Raymond, 2013 #288;Zheng, 2016 #16}. Special attention 1746 1747 should be given to these two hereditary syndromes, because for them there are well-1748 established screening guidelines available {Stoffel, 2015 #367;Daly, 2017 #368;Kratz, 2017 1749 #370;Ballinger, 2017 #369;Gupta, 2017 #366}. Up to 13% of all adrenal lesions in patients 1750 with MEN1 represent adrenal cancer (22084155). Cases of ACC have been reported in patients with Beckwith-Wiedemann syndrome (children), Familial Adenomatous Polyposis 1751 1752 (APC) and Carney Complex {Petr, 2016 #34}.

1753 Germline genetic testing for ACC patients should primarily be considered for the genes 1754 related to Li-Fraumeni syndrome and Lynch syndrome. ACC is an integral part of Li-1755 Fraumeni syndrome and when considering germline genetic testing, it is important to keep in 1756 mind that at least 20% of germline TP53 pathogenic variants occur as de novo mutations in 1757 the absence of any family history. Lynch syndrome is present in the same fraction of ACC 1758 patients as in colorectal cancer patients (3-5%), where general screening for Lynch syndrome is recommended {Stoffel, 2015 #367;Stoffel, 2015 #372}. Both, Li-Fraumeni 1759 1760 syndrome and Lynch syndrome have well established surveillance guidelines for carriers of 1761 pathogenic variants {Stoffel, 2015 #367;Daly, 2017 #368;Kratz, 2017 #370;Ballinger, 2017 1762 #369;Gupta, 2017 #366}. Evaluation for Lynch syndrome can be initiated by immunohistochemistry for MSH2, MLH1, PMS2, MSH6 and microsatellite instability testing, 1763 1764 or direct genetic germline analysis of MSH2, MLH1, PMS2, MSH6 and EPCAM. Genetic 1765 diagnosis of Li-Fraumeni syndrome is usually done by germline analysis for variants in TP53.

1766 For other syndromes (depending on family history and clinical suspicion) we refer to other 1767 sources {Petr, 2016 #34;Else, 2012 #290}.

Although not the topic of this guideline, all children with a diagnosis of ACC should undergo a systematic search of germline *TP53* pathogenic variants, because 50-90% of ACC in children are related to germline pathogenic *TP53* variants {McDonnell, 2003 #291;Custodio, 2013 #293;Wasserman, 2015 #292}

1772 1773

1776

1774**R.11.2.** The panel does not recommend for or against genetic tumor testing for1775somatic alterations.

1777 <u>Reasoning</u>

1778 While the panel recognizes that there is great hope that testing for somatic mutations and 1779 other markers in cancers general may allow tailoring of therapy and personalized 1780 approaches for therapy, for ACCs this approach is not yet established in routine clinical 1781 practice. Therefore, molecular testing should be offered within the framework of structured 1782 and systematic research projects.

- 1783
- 1784 1785

1786 **5.12. Pregnancy and ACC**

1787 1788

1789R.12.1. When an adrenal mass suspected to be an ACC is diagnosed during1790pregnancy, we recommend prompt surgical resection regardless of pregnancy1791trimester.

1792 1793 Reasoning

1794 Considering the poor prognosis of ACC and the importance of a prompt and complete 1795 surgical removal for prognosis, adrenal surgery should be pursued independent of the term 1796 of the pregnancy {Eschler, 2015 #129}. Preterm delivery (especially in the third trimester) 1797 and pregnancy loss are obvious risks when surgery is performed. Therefore, the patient and 1798 their family, obstetric providers and the ACC care team must engage in an informed 1799 discussion considering disease prognosis and the risk to the mother and fetus as related to 1800 the underlying disease and interventional procedures. A shared decision-making after 1801 discussion of all options is imperative.

1802 1803

1804**R.12.2.** Patients should be informed on pregnancy-related concerns specific to the1805current or past diagnosis of ACC.

- 1806
- 1807 <u>Reasoning</u>

1808 No evidence is available regarding how long patients should wait after the treatment of an1809 ACC before they can safely consider pregnancy.

1810 Importantly, the main concern is the poor prognosis of the malignant tumor and the potential 1811 that pregnancy could be a negative prognostic factor, possibly increasing the risk of 1812 recurrence. There is limited evidence that ACC occurring during pregnancy or in the 1813 postpartum period is associated with a worse prognosis than in non-pregnant women

- 1814 {Abiven-Lepage, 2010 #315}. The hypothesis that pregnancy could favor the development of1815 a more aggressive variant of ACC was raised.
- 1816 Due to the extreme paucity of information about this issue, it seems prudent to relay the 1817 information to the patient that there is a substantial risk of disease recurrence in the first 1818 years following the diagnosis of ACC.
- 1819 Since ACC may express estrogen receptors and there are preclinical data showing that 1820 estrogen may facilitate tumor development and progression through cross-talk with the IGF 1821 pathway {Sirianni, 2012 #317}, contraceptive measures other than estrogen-containing 1822 preparations are preferred.
- 1823
- 1824

1825 **R.12.3.** We recommend avoiding pregnancy while being on mitotane treatment.

1826 1827 Re

Reasoning 1828 The main concern with mitotane therapy is the potential of teratogenic effects, due to the 1829 suspicion that the drug may cross the placenta and cause adrenolytic activity on the human 1830 fetus. However, there are only few case reports of pregnancies when on mitotane therapy 1831 {Tripto-Shkolnik, 2013 #130}. Therefore, it is impossible to draw definitive conclusions about the safety of mitotane treatment or its associated risks. Woman treated with mitotane should 1832 1833 be informed about these risks, and ensure effective contraception to avoid pregnancy. 1834 Moreover, when mitotane treatment is discontinued, it seems wise to ensure undetectable 1835 mitotane plasma levels before considering pregnancy {de Corbiere, 2015 #318}, which might 1836 take 3-12 months. In case a patient becomes pregnant while on mitotane therapy, the 1837 uncertainty regarding risks of mitotane for the fetus should be discussed. In case the patient 1838 wishes to continue pregnancy mitotane therapy should be withheld.

- 1839 1840
- 1841
- 1842

1843

1844

43 **<u>6. Future directions and recommended research</u>**

1845 Due to the fact that the evidence for most of the recommendations provided in these 1846 guidelines is weak or even very weak, there are no doubts that major efforts are needed to 1847 improve diagnosis, treatment, and quality of life for patients with ACC.

Among many important research questions, we selected ten topics as particularly important.All of them can only be addressed in an international collaborative interdisciplinary manner.

1850 1) Clinical response to the best available therapy (i.e. EDP + mitotane) for advanced ACC is 1851 very limited with an objective response rate of less than 25%. Therefore, we underliably 1852 lack efficient drugs for treating this disease. Thus, identifying new therapeutic targets and 1853 options is a high priority. Here is a comprehensive but by far not complete list of 1854 emerging therapies: internal radionuclide therapy, such as metomidate-based therapies; 1855 drugs targeting the following pathways or targets: Wnt/beta-catenin; CDKN2A / TP53 / 1856 RB; IGF2 / mTOR; telomeres; drugs targeting histone modifications. In general, a 1857 combined approach seems to be reasonable. There is a growing notion that individual patients and individual tumors might respond better to certain therapies, depending on 1858 1859 their molecular landscape. Therefore, studies focusing on subgroup classification and 1860 identification are important. Due to the mitotane-associated pharmacological issues (e.g.

1861 CYP3A4 induction), it might be reasonable to test experimental drugs in mitotane-naïve 1862 patients within clinical studies.

- 1863 2) Immunotherapy is the latest revolution in cancer therapy, however preliminary data with
 1864 single immune check point inhibitors showed a modest activity in ACC patients. Molecular
 1865 and oncogenic pathways either in tumor cells or tumor microenvironment that can impair
 1866 induction or execution of a local antitumor immune response should be carefully studied in
 1867 ACC.
- 3) Since currently available systemic therapies have limited efficacy, but a subgroup of patient is destined to obtain a consistent benefit from them, the identification of predictive markers of efficacy (either clinical or molecular) of standard treatments is of paramount importance in order to spare toxic regimens to patients not destined to obtain a disease response.
- 4) With regards to improvement of surgery for ACC, standardization of procedures (e.g.
 laparoscopic vs. open surgery, lymph node dissection) should be promoted and tested in
 clinical trials.
- 5) The high recurrence rate in the majority of patients even after complete resection calls for improvement of adjuvant therapy. There are significant gaps in our understanding, which patients might truly benefit from the different adjuvant therapies and prospective trials are urgently needed. The ongoing ADIUVO trial will hopefully provide important information for low/intermediate risk patients, but a trial in high-risk patients (e.g. mitotane vs. mitotane + cisplatin + etoposide) is equally important.
- 6) Despite extensive efforts, the mechanism of action and pharmacodynamics of mitotane remain poorly understood {Hescot, 2015 #338;Hescot, 2013 #339;Sbiera, 2015 #337;Hescot, 2017 #340}. In addition, mitotane is a strong inducer of xenobiotics metabolism, probably negatively impacting subsequent and parallel therapies. Therefore, further understanding and improving the pharmacology and mechanism of action of mitotane with the goal of development of mitotane related drugs that do not share the negative adverse-effects would be a significant goal.
- 7) Translational research with the goal of rational treatment stratification should be promoted. 1889 1890 Recent molecular classifications, identifying distinct molecular subtypes with different 1891 outcomes, should be tested prospectively. These markers could provide a cornerstone for 1892 stratifying treatment strategies. This would mean that some patients of the 'better 1893 outcome' molecular group might benefit from forgoing any adjuvant therapy. Reversely, 1894 patients in the "poor outcome" molecular group could be included in a randomized trial 1895 testing mitotane + cytotoxic drugs as an adjuvant therapy. In addition, it will be important to define differences in pharmacogenomics or tumor genomics that define exceptional 1896 responders to mitotane and/or EDP. This data can fuel further sub-stratification of ACC 1897 1898 patients for certain therapies.
- 1899 8) In addition to improving treatment, other future research directions may include the use of 1900 artificial intelligence in diagnostic work-up of adrenal tumors and the improvement of screening and follow-up procedures using non-invasive techniques such as urine or 1901 1902 serum steroid metabolomics {Arlt, 2011 #48;Kerkhofs, 2015 #40;Taylor, 2017 #341;Hines, 1903 2017 #342} or 'liquid biopsies' with circulating tumor cells {Pinzani, 2013 #343}, circulating miRNAs {Chabre, 2013 #344;Szabo, 2014 #345;Perge, 2017 #346}, or circulating cell-free 1904 1905 tumor DNA {Creemers, 2017 #347;Garinet, 2018 #348} for early diagnosis or detection of 1906 recurrence.
- 1907 9) In the long term, a better understanding of the pathogenesis of ACC is needed to pave the1908 way for future progress. Therefore, basic research efforts have to continue. Preclinical

1909 models are needed, to test new treatments, including additional new cell lines, tumor organoids, and new animal models. Mechanisms of tumorigenesis, tumor evolution 1910 (genetic heterogeneity, clonal evolution) and further definition of known and future 1911 therapeutic targets should be encouraged. 1912

- 1913 10) No studies so far have revealed the wishes and experiences of patients. Given the poor prognosis and the toxic therapies, there is a definite need for 'Patient Related Outcomes'. 1914 1915 PRO's should be measured (PROM's) and incorporated in our strategy for value based 1916 cure and care.
- 1917

1918 In general, it is our common task to overcome the major limitation in ACC research - the 1919 rarity of this disease. Therefore, beyond proofs of concept requiring few patients, clinical 1920 trials can only be performed if a large number of centers gather multicenter studies. This 1921 underscores the critical role of adrenal research networks, such as ENSAT or A5, to 1922 coordinate these efforts. Ideally a limited number of large prospective trials should continuously be ongoing, in order to allow for sufficient patient recruitment. In the same 1923 1924 context we envision that at least one reference center in every country will be established to 1925 provide multidisciplinary expertise for this rare disease to all patients.

1926

1927 Altogether, owing to its rarity and its severity, ACC should continue to mobilize researchers, physicians and patients in a coordinated engaged effort. 1928 1929

1930

1931 1932

1933 Appendices

- 1934 Appendix 1: Question 1: Pathology - what is needed to diagnose an ACC? Summary of included studies (1a: distinguishing adrenal from non-adrenal tumors; 1b: 1935 1936 distinguishing benign from malignant behavior in adrenal tumors)
- 1937 Appendix 2: Question 2: Which are the best prognostic markers in ACC? Summary of 1938 included studies 1939
 - Appendix 3: Question 2: Prognostic factors in ACC overview of studies markers
- 1940 Appendix 4: Question 3: Is adjuvant therapy able to prevent recurrent disease or 1941 reduce mortality after radical resection? Summary of included studies (3a: Adjuvant mitotane after surgery; 3b: Adjuvant radiotherapy after surgical resection) 1942
 - Appendix 5: Evidence tables Question 3 (adjuvant therapy)
 - Appendix 6: Question 4: What is the best treatment option for macroscopically incompletely resected, recurrent or metastatic disease? Summary of included studies
- 1946 • Appendix 7: Summary of relevant drug interactions with mitotane
- Appendix 8: Comments to this Guidelines by invited reviewers and members of the 1947 European Society of Endocrinology (ESE) and the European Network for the Study of 1948 1949 Adrenal Tumors (ENSAT), representatives of associated societies of ESE, and 1950 patient representatives
- 1951 1952

1943

1944

1945

1953 Fundina

1954 This guideline was sponsored by the European Society of Endocrinology with support by the European Network for the Study of Adrenal Tumors. 1955 1956

Acknowledgements 1957

The authors of the guideline would like to thank and acknowledge Mouhammed Habra, 1958 1959 Electron Kebebev, and Britt Skogseid for their expert review and additional members of the European Society of Endocrinology, the European Network for the Study of Adrenal Tumors 1960

1961 or representatives of national endocrine societies for valuable and critical comments. In 1962 addition, we thank 3 patient representatives who provided valuable feedback for the 1963 guideline. The comments of the reviewers as well as the authors' responses are available as Appendix 8. Furthermore, we thank John Newell-Price for very helpful English proof-reading. 1964 1965 Finally we would like to thank Annemarie Venemans for her support in the systematic 1966 literature search. 1967 1968 1969 Declaration of potential conflict of interests (in the last 5 years) 1970 1971 Guillaume Assié 1972 • Speakers fee / travel support for congresses from: HRA Pharma (2016); Ipsen Pharma (2013, 2014); Novartis (2012, 2013, 2014, 2015, 2016) 1973 1974 Eric Baudin 1975 1976 Speakers fee: HRA Pharma • 1977 Research support by HRA Pharma • 1978 1979 Alfredo Berruti 1980 Member to remunerated Advisory Boards of Astellas, Sanofi, Janssen, Merck Sharp • 1981 and Dome, Novartis, Ipsen 1982 • Speakers fee / travel support for congresses from: Astellas, Sanofi, Janssen, 1983 Novartis, Ipsen 1984 Research support by Janssen (Phase II trial of Abiraterone in the management of • Cushing Syndrome induced by Adrennocortical Carcinoma; 2016); Sanofi: Phase II 1985 trial of Cabazitaxel as second line treatment in the treatment of patients with 1986 1987 advanced Adrenocortical Carcinoma; 2014) 1988 1989 Martin Fassnacht 1990 Advisory board member: of HRA Pharma (2015; not remunerated); Atterocor (2013); 1991 Astellas Pharma (2012) 1992 Speakers fee / travel support for congresses from: HRA Pharma (2013); Ipsen • 1993 Pharma (2011, 2012) 1994 1995 Harm Haak 1996 Research support by HRA Pharma (2016) 1997 1998 Massimo Terzolo -• Advisory Board member of HRA Pharma (2013; not remunerated), Atterocor-Millendo 1999 2000 (2013 - 2015)2001 Research support by HRA Pharma (2016) Speaker fee/travel support from HRA Pharma (2014, 2015) 2002 2003 2004 The other authors declare no conflict of interest. 2005 2006

2007 **References**

2008 1 Kebebew E, Reiff E, Duh QY, Clark OH & McMillan A. Extent of disease at
2009 presentation and outcome for adrenocortical carcinoma: have we made progress? World
2010 Journal of Surgery 2006 30 872–878. (https://doi.org/10.1007/s00268-005-0329-x)

2011 2 Kerkhofs TM, Verhoeven RH, Van der Zwan JM, Dieleman J, Kerstens MN, Links TP, 2012 Van de Poll-Franse LV & Haak HR. Adrenocortical carcinoma: a population-based study on 2013 incidence and survival in the Netherlands since 1993. European Journal of Cancer 2013 49 2014 2579–2586. (https://doi.org/10.1016/j.ejca.2013.02.034)

Berruti A, Baudin E, Gelderblom H, Haak HR, Porpiglia F, Fassnacht M &
Pentheroudakis G. Adrenal cancer: ESMO Clinical Practice Guidelines for diagnosis,
treatment and follow-up. Annals of Oncology 2012 23 131–138.

2018 4 Petr EJ & Else T. Genetic predisposition to endocrine tumors: diagnosis, surveillance
2019 and challenges in care. Seminars in Oncology 2016 43 582–590. (https://doi.org/10.1053/j.
2020 seminoncol.2016.08.007)

de Reynies A, Assie G, Rickman DS, Tissier F, Groussin L, Rene- Corail F, Dousset
B, Bertagna X, Clauser E & Bertherat J. Gene expression profiling reveals a new
classification of adrenocortical tumors and identifies molecular predictors of malignancy and
survival. Journal of Clinical Oncology 2009 27 1108–1115. (https://doi.
org/10.1200/JCO.2008.18.5678)

Fragoso MC, Almeida MQ, Mazzuco TL, Mariani BM, Brito LP, Goncalves TC,
Alencar GA, Lima L de O, Faria AM, Bourdeau I et al. Combined expression of BUB1B,
DLGAP5, and PINK1 as predictors of poor outcome in adrenocortical tumors: validation in a
Brazilian cohort of adult and pediatric patients. European Journal of Endocrinology 2012 166
61–67. (https://doi.org/10.1530/EJE-11-0806)

7 Ronchi CL, Sbiera S, Leich E, Henzel K, Rosenwald A, Allolio B & Fassnacht M. Single
nucleotide polymorphism array profiling of adrenocortical tumors – evidence for an adenoma
carcinoma sequence? PLoS ONE 2013 8 e73959. (https://doi.org/10.1371/
journal.pone.0073959)

2035 8 Jouinot A, Assie G, Libe R, Fassnacht M, Papathomas T, Barreau O, de la Villeon B,
2036 Faillot S, Hamzaoui N, Neou M et al. DNA methylation is an independent prognostic marker
2037 of survival in adrenocortical cancer. Journal of Clinical Endocrinology and Metabolism 2017
2038 102 923–932.

- Assie G, Letouze E, Fassnacht M, Jouinot A, Luscap W, Barreau O, Omeiri H,
 Rodriguez S, Perlemoine K, Rene-Corail F et al. Integrated genomic characterization of
 adrenocortical carcinoma. Nature Genetics 2014 46 607–612.
- 2042 (https://doi.org/10.1038/ng.2953)
- 2043 10 Juhlin CC, Goh G, Healy JM, Fonseca AL, Scholl UI, Stenman A, Kunstman JW,
 2044 Brown TC, Overton JD, Mane SM et al. Whole-exome sequencing characterizes the
 2045 landscape of somatic mutations and copy number alterations in adrenocortical carcinoma.
 2046 Journal of Clinical Endocrinology and Metabolism 2015 100 E493–E502. (https://
 2047 doi.org/10.1210/jc.2014-3282)
- 2048 11 Zheng S, Cherniack AD, Dewal N, Moffitt RA, Danilova L, Murray BA, Lerario AM,
 2049 Else T, Knijnenburg TA, Ciriello G et al. Comprehensive pan-genomic characterization of
 2050 adrenocortical carcinoma. Cancer Cell 2016 29 723–736.
- 2051 (https://doi.org/10.1016/j.ccell.2016.04.002)

2052 12 Assie G, Jouinot A & Bertherat J. The 'omics' of adrenocortical tumours for
2053 personalized medicine. Nature Reviews Endocrinology 2014 10 215–228.
2054 (https://doi.org/10.1038/nrendo.2013.272)

2055 13 Else T, Kim AC, Sabolch A, Raymond VM, Kandathil A, Caoili EM, Jolly S, Miller BS,
2056 Giordano TJ & Hammer GD. Adrenocortical carcinoma. Endocrine Reviews 2014 35 282–
2057 326. (https://doi. org/10.1210/er.2013-1029)

2058 14 Faillot S & Assie G. ENDOCRINE TUMOURS: The genomics of adrenocortical
2059 tumors. European Journal of Endocrinology 2016 174 R249–R265.
2060 (https://doi.org/10.1530/EJE-15-1118)

- 15 Terzolo M, Ali A, Osella G & Mazza E. Prevalence of adrenal carcinoma among
 incidentally discovered adrenal masses. A retrospective study from 1989 to 1994. Gruppo
 Piemontese Incidentalomi Surrenalici. Archives of Surgery 1997 132 914–919.
 (https://doi.org/10.1001/archsurg.1997.01430320116020)
- 2065 16 Cawood TJ, Hunt PJ, O'Shea D, Cole D & Soule S. Recommended evaluation of
 2066 adrenal incidentalomas is costly, has high false-positive rates and confers a risk of fatal
 2067 cancer that is similar to the risk of the adrenal lesion becoming malignant; time for a rethink?
 2068 European Journal of Endocrinology 2009 161 513–527. (https://doi.org/10.1530/ EJE-092069 0234)
- 2070 17 Fassnacht M, Arlt W, Bancos I, Dralle H, Newell-Price J, Sahdev A, Tabarin A,
 2071 Terzolo M, Tsagarakis S & Dekkers OM. Management of adrenal incidentalomas: European
 2072 Society of Endocrinology Clinical Practice Guideline in collaboration with the European
 2073 Network for the Study of Adrenal Tumors. European Journal of Endocrinology 2016 175 G1–
 2074 G34. (https://doi.org/10.1530/EJE-16-0467)
- 2075 18 Seccia TM, Fassina A, Nussdorfer GG, Pessina AC & Rossi GP. Aldosterone2076 producing adrenocortical carcinoma: an unusual cause of Conn's syndrome with an ominous
 2077 clinical course. Endocrine-Related Cancer 2005 12 149–159.
 2078 (https://doi.org/10.1677/org.1.00267)
- 2078 (https://doi.org/10.1677/erc.1.00867)
- 2079 19 Fassnacht M, Libe R, Kroiss M & Allolio B. Adrenocortical carcinoma: a clinician's
 2080 update. Nature Reviews Endocrinology 2011 7 323–335.
 2081 (https://doi.org/10.1038/nrendo.2010.235)
- 2082 20 Berruti A, Fassnacht M, Haak H, Else T, Baudin E, Sperone P, Kroiss M, Kerkhofs T,
 2083 Williams AR, Ardito A et al. Prognostic role of overt hypercortisolism in completely operated
 2084 patients with adrenocortical cancer. European Urology 2014 65 832–838.
 2085 (https://doi.org/10.1016/j. eururo.2013.11.006)
- 2086 21 Kerkhofs TM, Ettaieb MH, Hermsen IG & Haak HR. Developing treatment for
 2087 adrenocortical carcinoma. Endocrine-Related Cancer 2015 22 R325–R338.
 2088 (https://doi.org/10.1530/ERC-15-0318)
- 2089 22 Fassnacht M, Kroiss M & Allolio B. Update in adrenocortical carcinoma. Journal of
 2090 Clinical Endocrinology and Metabolism 2013 98 4551–4564. (https://doi.org/10.1210/jc.20132091 3020)
- 209223Fassnacht M & Allolio B. Clinical management of adrenocortical carcinoma. Best2093Practice and Research Clinical Endocrinology and Metabolism 2009 23 273–289.
- 2094 24 Johanssen S, Hahner S, Saeger W, Quinkler M, Beuschlein F, Dralle H, Haaf M,
 2095 Kroiss M, Jurowich C, Langer P et al. Deficits in the management of patients with

adrenocortical carcinoma in germany. Deutsches Arzteblatt International 2010 107 U885–
U889.

2098 25 Icard P, Goudet P, Charpenay C, Andreassian B, Carnaille B, Chapuis Y, Cougard P,
2099 Henry JF & Proye C. Adrenocortical carcinomas: surgical trends and results of a 253-patient
2100 series from the French Association of Endocrine Surgeons study group. World Journal of
2101 Surgery 2001 25 891–897. (https://doi.org/10.1007/s00268- 001-0047-y)

26 Bilimoria KY, Shen WT, Elaraj D, Bentrem DJ, Winchester DJ, Kebebew E &
Sturgeon C. Adrenocortical carcinoma in the United States: treatment utilization and
prognostic factors. Cancer 2008 113 3130–3136. (https://doi.org/10.1002/cncr.23886)

2105 27 Sturgeon C, Shen WT, Clark OH, Duh QY & Kebebew E. Risk assessment in 457
2106 adrenal cortical carcinomas: how much does tumor size predict the likelihood of malignancy?
2107 Journal of the American College of Surgeons 2006 202 423–430. (https://doi.
2108 org/10.1016/j.jamcollsurg.2005.11.005)

28 Fassnacht M, Johanssen S, Fenske W, Weismann D, Agha A, Beuschlein F, Fuhrer
D, Jurowich C, Quinkler M, Petersenn S et al. Improved survival in patients with stage II
adrenocortical carcinoma followed up prospectively by specialized centers. Journal of Clinical
Endocrinology and Metabolism 2010 95 4925–4932.

29 Fassnacht M, Johanssen S, Quinkler M, Bucsky P, Willenberg HS, Beuschlein F,
Terzolo M, Mueller HH, Hahner S & Allolio B. Limited prognostic value of the 2004
International Union Against Cancer staging classification for adrenocortical carcinoma:
proposal for a Revised TNM Classification. Cancer 2009 115 243–250. (https://doi.
org/10.1002/cncr.24030)

30 Fassnacht M, Terzolo M, Allolio B, Baudin E, Haak H, Berruti A, Welin S, SchadeBrittinger C, Lacroix A, Jarzab B et al. Combination chemotherapy in advanced
adrenocortical carcinoma. New England Journal of Medicine 2012 366 2189–2197.
(https://doi.org/10.1056/ NEJMoa1200966)

31 Bollerslev J, Rejnmark L, Marcocci C, Shoback DM, Sitges-Serra A, van Biesen W &
2123 Dekkers OM. European Society of Endocrinology Clinical Guideline: treatment of chronic
2124 hypoparathyroidism in adults. European Journal of Endocrinology 2015 173 G1–G20.
2125 (https:// doi.org/10.1530/EJE-15-0628)

Andrews J, Guyatt G, Oxman AD, Alderson P, Dahm P, Falck-Ytter Y, Nasser M,
Meerpohl J, Post PN, Kunz R et al. GRADE guidelines: 14. Going from evidence to
recommendations: the significance and presentation of recommendations. Journal of Clinical
Epidemiology 2013 66 719–725. (https://doi.org/10.1016/j.jclinepi.2012.03.013)

Andrews JC, Schunemann HJ, Oxman AD, Pottie K, Meerpohl JJ, Coello PA, Rind D,
Montori VM, Brito JP, Norris S et al. GRADE guidelines: 15. Going from evidence to
recommendation- determinants of a recommendation's direction and strength. Journal of
Clinical Epidemiology 2013 66 726–735. (https://doi.org/10.1016/j. jclinepi.2013.02.003)

34 Guyatt GH, Schunemann HJ, Djulbegovic B & Akl EA. Guideline panels should not
GRADE good practice statements. Journal of Clinical Epidemiology 2015 68 597–600.
(https://doi.org/10.1016/j. jclinepi.2014.12.011)

2137 35 Vanbrabant T, Fassnacht M, Assie G & Dekkers OM. Influence of hormonal functional
2138 status on survival in adrenocortical carcinoma: systematic review and meta-analysis.
2139 European Journal of Endocrinology 2018.

- Bertherat J, Coste J & Bertagna X. Adjuvant mitotane in adrenocortical carcinoma.
 New England Journal of Medicine 2007 357 1256–1257; author reply 1259.
- 2142 (https://doi.org/10.1056/ NEJMc076267)

2143 37 Blanes A & Diaz-Cano SJ. Histologic criteria for adrenocortical proliferative lesions –
2144 value of mitotic figure variability. American Journal of Clinical Pathology 2007 127 398–408.
2145 (https://doi. org/10.1309/MCGUQ3R4A4WWN3LB)

2146 38 Creemers SG, van Koetsveld PM, van Kemenade FJ, Papathomas TG, Franssen GJ,
2147 Dogan F, Eekhoff EM, van der Valk P, de Herder WW, Janssen JA et al. Methylation of IGF2
2148 regulatory regions to diagnose adrenocortical carcinomas. Endocrine-Related Cancer 2016
2149 23 727–737. (https://doi.org/10.1530/ERC-16-0266)

2150 39 Erickson LA, Jin L, Sebo TJ, Lohse C, Pankratz VS, Kendrick ML, van Heerden JA,
2151 Thompson GB, Grant CS & Lloyd RV. Pathologic features and expression of insulin-like
2152 growth factor-2 in adrenocortical neoplasms. Endocrine Pathology 2001 12 429–435.
2153 (https://doi. org/10.1385/EP:12:4:429)

40 Arola J, Liu J, Heikkila P, Ilvesmaki V, Salmenkivi K, Voutilainen R & Kahri AI.
Expression of inhibin alpha in adrenocortical tumours reflects the hormonal status of the
neoplasm. Journal of Endocrinology

2157 2000 165 223–229. (https://doi.org/10.1677/joe.0.1650223)

2158 41 Aubert S, Wacrenier A, Leroy X, Devos P, Carnaille B, Proye C, Wemeau JL, Lecomte-

2159 Houcke M & Leteurtre E. Weiss system revisited: a clinicopathologic and

2160 immunohistochemical study of 49 adrenocortical tumors. American Journal of Surgical

2161 Pathology 2002 26 1612–1619. (https://doi.org/10.1097/00000478-200212000-00009)

42 Busam KJ, Iversen K, Coplan KA, Old LJ, Stockert E, Chen YT, McGregor D &
Jungbluth A. Immunoreactivity for A103, an antibody to melan-A (Mart-1), in adrenocortical
and other steroid tumors. American Journal of Surgical Pathology 1998 22 57–63.
(https://doi. org/10.1097/00000478-199801000-00007)

43 Kamio T, Shigematsu K, Sou H, Kawai K & Tsuchiyama H. Immunohistochemical
expression of epidermal growth factor receptors in human adrenocortical carcinoma. Human
Pathology 1990 21 277–282. (https://doi.org/10.1016/0046-8177(90)90227-V)

2169 44 Komminoth P, Roth J, Schroder S, Saremaslani P & Heitz PU. Overlapping

2170 expression of immunohistochemical markers and synaptophysin mRNA in

2171 pheochromocytomas and adrenocortical carcinomas. Implications for the differential

2172 diagnosis of adrenal gland tumors. Laboratory Investigation 1995 72 424–431.

45 Pan CC, Chen PCH, Tsay SH & Ho DMT. Differential immunoprofiles of
hepatocellular carcinoma, renal cell carcinoma, and adrenocortical carcinoma: a systemic
immunohistochemical survey using tissue array technique. Applied Immunohistochemistry
and Molecular Morphology 2005 13 347–352. (https://doi.org/10.1097/01.

2177 pai.0000146525.72531.19)

46 Rubin B, Regazzo D, Redaelli M, Mucignat C, Citton M, Iacobone M, Scaroni C,
Betterle C, Mantero F, Fassina A et al. Investigation of N-cadherin/beta-catenin expression in
adrenocortical tumors. Tumour Biology 2016 37 13545–13555. (https://doi.org/10.1007/
s13277-016-5257-x)

2182 47 Sbiera S, Schmull S, Assie G, Voelker HU, Kraus L, Beyer M, Ragazzon B,
2183 Beuschlein F, Willenberg HS, Hahner S et al. High diagnostic and prognostic value of

- steroidogenic factor-1 expression in adrenal tumors. Journal of Clinical Endocrinology and
 Metabolism 2010 95 E161–E171.
- 48 Stojadinovic A, Brennan MF, Hoos A, Omeroglu A, Leung DH, Dudas ME, Nissan A,
 Cordon-Cardo C & Ghossein RA. Adrenocortical adenoma and carcinoma: histopathological
 and molecular comparative analysis. Modern Pathology 2003 16 742–751.
 (https://doi.org/10.1097/01.MP.0000081730.72305.81)
- 49 Volante M, Sperone P, Bollito E, Frangipane E, Rosas R, Daffara F, Terzolo M,
 Berruti A & Papotti M. Matrix metalloproteinase type 2 expression in malignant adrenocortical
 tumors: Diagnostic and prognostic significance in a series of 50 adrenocortical carcinomas.
 Modern Pathology 2006 19 1563–1569. (https://doi.org/10.1038/ modpathol.3800683)
- Wajchenberg BL, Albergaria Pereira MA, Medonca BB, Latronico AC, Campos
 Carneiro P, Alves VA, Zerbini MC, Liberman B, Carlos Gomes G & Kirschner MA.
 Adrenocortical carcinoma: clinical and laboratory observations. Cancer 2000 88 711–736.
 (https:// doi.org/10.1002/(SICI)1097-0142(20000215)88:4<711::AID- CNCR1>3.0.CO;2-W)
- 2198 51 Wang C, Sun Y, Wu H, Zhao D & Chen J. Distinguishing adrenal cortical carcinomas
 2199 and adenomas: a study of clinicopathological features and biomarkers. Histopathology 2014
 2200 64 567–576. (https:// doi.org/10.1111/his.12283)
- 2201 52 Zhang HY, Bu H, Chen HJ, Wei B, Liu WP, Guo J, Li FY, Liao DY, Tang Y & Zhang Z.
 2202 Comparison of immunohistochemical markers in the differential diagnosis of adrenocortical
 2203 tumors immunohistochemical analysis of adrenocortical tumors. Applied
 2204 Immunohistochemistry and Molecular Morphology 2008 16 32–39.
- 53 Kovach AE, Nucera C, Lam QT, Nguyen A, Dias-Santagata D & Sadow PM. Genomic
 and immunohistochemical analysis in human adrenal cortical neoplasia reveal beta-catenin
 mutations as potential prognostic biomarker. Discoveries 2015 3 e40.
- Amini N, Margonis GA, Kim Y, Tran TB, Postlewait LM, Maithel SK, Wang TS, Evans
 DB, Hatzaras I, Shenoy R et al. Curative resection of adrenocortical carcinoma: rates and
 patterns of postoperative recurrence. Annals of Surgical Oncology 2016 23 126–133.
 (https://doi. org/10.1245/s10434-015-4810-y)
- 55 Asare EA, Wang TS, Winchester DP, Mallin K, Kebebew E & Sturgeon C. A novel
 staging system for adrenocortical carcinoma better predicts survival in patients with stage I/II
 disease. Surgery 2014 156 1378–1386. (https://doi.org/10.1016/j. surg.2014.08.018)
- S6 Assie G, Antoni G, Tissier F, Caillou B, Abiven G, Gicquel C, Leboulleux S, Travagli
 JP, Dromain C, Bertagna X et al. Prognostic parameters of metastatic adrenocortical
 carcinoma. Journal of Clinical Endocrinology and Metabolism 2007 92 148–154.
- Ayala-Ramirez M, Jasim S, Feng L, Ejaz S, Deniz F, Busaidy N, Waguespack SG,
 Naing A, Sircar K, Wood CG et al. Adrenocortical carcinoma: clinical outcomes and
 prognosis of 330 patients at a tertiary care center. European Journal of Endocrinology 2013
 169 891–899. (<u>https://doi.org/10.1530/EJE-13-0519</u>)
- 58 Beuschlein F, Weigel J, Saeger W, Kroiss M, Wild V, Daffara F, Libe R, Ardito A,
 Ghuzlan AA, Quinkler M et al. Major prognostic role of Ki67 in localized adrenocortical
 carcinoma after complete resection. Journal of Clinical Endocrinology and Metabolism 2015
 100 841–849. (https://doi.org/10.1210/jc.2014-3182)
- 2226 59 Canter DJ, Mallin K, Uzzo RG, Egleston BL, Simhan J, Walton J, Smaldone MC,
 2227 Master VA, Bratslaysky G & Kutikov A. Association of tumor size with metastatic potential

- and survival in patients with adrenocortical carcinoma: an analysis of the National Cancer
 Database. Canadian Journal of Urology 2013 20 6915–6921.
- 2230 60 Duregon E, Cappellesso R, Maffeis V, Zaggia B, Ventura L, Berruti A, Terzolo M,
 2231 Fassina A, Volante M & Papotti M. Validation of the prognostic role of the 'Helsinki Score' in
 2232 225 cases of adrenocortical carcinoma. Human Pathology 2017 62 1–7.
 2233 (https://doi.org/10.1016/j. humpath.2016.09.035)
- Erdogan I, Deutschbein T, Jurowich C, Kroiss M, Ronchi C, Quinkler M, Waldmann J,
 Willenberg HS, Beuschlein F, Fottner C et al. The role of surgery in the management of
 recurrent adrenocortical carcinoma. Journal of Clinical Endocrinology and Metabolism 2013
 98 181–191.
- Ettaieb MH, Duker JC, Feelders RA, Corssmit EP, Menke-van der Houven van Oordt
 CW, Timmers HJ, Kerstens MN, Wilmink JW, Zelissen PM, Havekes B et al. Synchronous vs
 metachronous metastases in adrenocortical carcinoma: an analysis of the dutch adrenal
 network. Hormones and Cancer 2016 7 336–344. (https://doi. org/10.1007/s12672-016-02705)
- Freire DS, Siqueira SAC, Zerbini MCN, Wajchenberg BL, Correa- Giannella ML,
 Lucon AM & Pereira MAA. Development and internal validation of an adrenal cortical
 carcinoma prognostic score for predicting the risk of metastasis and local recurrence. Clinical
 Endocrinology 2013 79 468–475. (https://doi.org/10.1111/ cen.12174)
- 64 Gicquel C, Bertagna X, Gaston V, Coste J, Louvel A, Baudin E, Bertherat J, Chapuis
 Y, Duclos JM, Schlumberger M et al. Molecular markers and long-term recurrences in a large
 cohort of patients with sporadic adrenocortical tumors. Cancer Research 2001 61 6762–
 6767.
- Glover AR, Zhao JT, Ip JC, Lee JC, Robinson BG, Gill AJ, Soon PS & Sidhu SB.
 Long noncoding RNA profiles of adrenocortical cancer can be used to predict recurrence.
 Endocrine-Related Cancer 2015 22 99–109. (https://doi.org/10.1530/ERC-14-0457)
- 66 Gonzalez RJ, Tamm EP, Ng C, Phan AT, Vassilopoulou-Sellin R, Perrier ND, Evans
 DB & Lee JE. Response to mitotane predicts outcome in patients with recurrent adrenal
 cortical carcinoma. Surgery 2007 142 867–874. (https://doi.org/10.1016/j. surg.2007.09.006)
- Kendrick ML, Curlee K, Lloyd R, Farley DR, Grant CS, Thompson GB, Rowland C,
 Young WF Jr, Van Heerden JA, Duh QY et al. Aldosterone-secreting adrenocortical
 carcinomas are associated with unique operative risks and outcomes. Surgery 2002 132
 1008–1012. (https://doi.org/10.1067/msy.2002.128476)
- Kim Y, Margonis GA, Prescott JD, Tran TB, Postlewait LM, Maithel SK, Wang TS,
 Evans DB, Hatzaras I, Shenoy R et al. Nomograms to predict recurrence-free and overall
 survival after curative resection of adrenocortical carcinoma. JAMA Surgery 2016 151 365–
 373. (https:// doi.org/10.1001/jamasurg.2015.4516)
- Kim Y, Margonis GA, Prescott JD, Tran TB, Postlewait LM, Maithel SK, Wang TS,
 Glenn JA, Hatzaras I, Shenoy R et al. Curative surgical resection of adrenocortical
 carcinoma: determining long-term outcome based on conditional disease-free probability.
 Annals of Surgery 2017 265 197–204. (https://doi.org/10.1097/ SLA.000000000001527)
- Libe R, Borget I, Ronchi CL, Zaggia B, Kroiss M, Kerkhofs T, Bertherat J, Volante M,
 Quinkler M, Chabre O et al. Prognostic factors in stage III-IV adrenocortical carcinomas
 (ACC): an European Network for the Study of Adrenal Tumor (ENSAT) study. Annals of
 Oncology 2015 26 2119–2125. (https://doi.org/10.1093/annonc/ mdv329)

2273 71 Livhits M, Li N, Yeh MW & Harari A. Surgery is associated with improved survival for
2274 adrenocortical cancer, even in metastatic disease. Surgery 2014 156 1531–1540; discussion
2275 1540–1531. (https:// doi.org/10.1016/j.surg.2014.08.047)

Lucon AM, Pereira MA, Mendonca BB, Zerbini MC, Saldanha LB & Arap S.
Adrenocortical tumors: results of treatment and study of Weiss's score as a prognostic factor.
Revista do Hospital das Clinicas da Faculdade de Medicina da Universidade de Sao Paulo
2002 57 251–256. (https://doi.org/10.1590/S0041-87812002000600002)

Margonis GA, Kim Y, Prescott JD, Tran TB, Postlewait LM, Maithel SK, Wang TS,
Evans DB, Hatzaras I, Shenoy R et al. Adrenocortical carcinoma: impact of surgical margin
status on long-term outcomes. Annals of Surgical Oncology 2016 23 134–141. (https://doi.
org/10.1245/s10434-015-4803-x)

Margonis GA, Kim Y, Tran TB, Postlewait LM, Maithel SK, Wang TS, Glenn JA,
Hatzaras I, Shenoy R, Phay JE et al. Outcomes after resection of cortisol-secreting
adrenocortical carcinoma. American Journal of Surgery 2016 211 1106–1113.
(https://doi.org/10.1016/j. amjsurg.2015.09.020)

Millis SZ, Ejadi S & Demeure MJ. Molecular profiling of refractory adrenocortical
cancers and predictive biomarkers to therapy. Biomarkers in Cancer 2015 7 69–76.
(https://doi.org/10.4137/BIC.S34292)

76 Paton BL, Novitsky YW, Zerey M, Harrell AG, Norton HJ, Asbun H, Kercher KW &
Heniford BT. Outcomes of adrenal cortical carcinoma in the United States. Surgery 2006 140
914–920; discussion 919–920. (https://doi.org/10.1016/j.surg.2006.07.035)

Pennanen M, Heiskanen I, Sane T, Remes S, Mustonen H, Haglund C & Arola J.
Helsinki score-a novel model for prediction of metastases in adrenocortical carcinomas.
Human Pathology 2015 46 404–410. (https://doi.org/10.1016/j.humpath.2014.11.015)

78 Schulick RD & Brennan MF. Long-term survival after complete resection and repeat
 resection in patients with adrenocortical carcinoma. Annals of Surgical Oncology 1999 6 719 726. (https://doi. org/10.1007/s10434-999-0719-7)

Tran TB, Postlewait LM, Maithel SK, Prescott JD, Wang TS, Glenn J, Phay JE,
Keplinger K, Fields RC, Jin LDX et al. Actual 10-year survivors following resection of
adrenocortical carcinoma. Journal of Surgical Oncology 2016 114 971–976.
(https://doi.org/10.1002/ jso.24439)

Xiao WJ, Zhu Y, Dai B, Zhang HL, Shi GH, Shen YJ, Zhu YP & Ye DW. Conditional
survival among patients with adrenal cortical carcinoma determined using a national
population-based surveillance, epidemiology, and end results registry. Oncotarget 2015 6
44955–44962.

Zini L, Capitanio U, Jeldres C, Lughezzani G, Sun M, Shariat SF, Isbarn H, Arjane P,
Widmer H, Perrotte P et al. External validation of a nomogram predicting mortality in patients
with adrenocortical carcinoma. BJU International 2009 104 1661–1667. (https://doi.
org/10.1111/j.1464-410X.2009.08660.x)

Ronchi CL, Sbiera S, Leich E, Tissier F, Steinhauer S, Deutschbein T, Fassnacht M &
Allolio B. Low SGK1 expression in human adrenocortical tumors is associated with ACTHindependent glucocorticoid secretion and poor prognosis. Journal of Clinical Endocrinology
and Metabolism 2012 97 E2251–E2260. (https://doi. org/10.1210/jc.2012-2669)

2316 83 Macfarlane DA. Cancer of the adrenal cortex; the natural history, prognosis and
2317 treatment in a study of fifty-five cases. Annals of the Royal College of Surgeons of England
2318 1958 23 155–186.

231984Sullivan M, Boileau M & Hodges CV. Adrenal cortical carcinoma. Journal of Urology23201978 120 660–665. (https://doi.org/10.1016/S0022-5347(17)57317-6)

2321 85 Lee JE, Berger DH, el-Naggar AK, Hickey RC, Vassilopoulou-Sellin R, Gagel RF,
2322 Burgess MA & Evans DB. Surgical management, DNA content, and patient survival in
2323 adrenal cortical carcinoma. Surgery 1995 118 1090–1098. (https://doi.org/10.1016/S00392324 6060(05)80119-9)

2325 86 DeLellis RA, Lloyd RV, Heitz PU & Eng C. World Health Organization classification of 2326 tumours. Pathology and Genetics of Tumours of Endocrine Organs 2004 136.

2327 87 Miller BS, Gauger PG, Hammer GD, Giordano TJ & Doherty GM. Proposal for
2328 modification of the ENSAT staging system for adrenocortical carcinoma using tumor grade.
2329 Langenbecks Archives of Surgery 2010 395 955–961. (https://doi.org/10.1007/s00423-0102330 0698-y)

2331 88 Lughezzani G, Sun M, Perrotte P, Jeldres C, Alasker A, Isbarn H, Budaus L, Shariat
2332 SF, Guazzoni G, Montorsi F et al. The European Network for the Study of Adrenal Tumors
2333 staging system is prognostically superior to the international union against cancer- staging
2334 system: a North American validation. European Journal of Cancer 2010 46 713–719.
2335 (https://doi.org/10.1016/j.ejca.2009.12.007)

2336 89 Lam AK. Update on adrenal tumours in 2017 World Health Organization (WHO) of
2337 endocrine tumours. Endocrine Pathology 2017 28 213–227. (https://doi.org/10.1007/s120222338 017-9484-5)

90 Berruti A, Grisanti S, Pulzer A, Claps M, Daffara F, Loli P, Mannelli M, Boscaro M,
Arvat E, Tiberio G et al. Long-term outcomes of adjuvant mitotane therapy in patients with
radically resected adrenocortical carcinoma. Journal of Clinical Endocrinology and
Metabolism 2017 102 1358–1365. (https://doi.org/10.1210/jc.2016-2894)

2343 91 Else T, Williams AR, Sabolch A, Jolly S, Miller BS & Hammer GD. Adjuvant therapies
2344 and patient and tumor characteristics associated with survival of adult patients with
2345 adrenocortical carcinoma. Journal of Clinical Endocrinology and Metabolism 2014 99 455–
2346 461. (https:// doi.org/10.1210/jc.2013-2856)

92 Grubbs EG, Callender GG, Xing Y, Perrier ND, Evans DB, Phan AT & Lee JE.
Recurrence of adrenal cortical carcinoma following resection: surgery alone can achieve
results equal to surgery plus mitotane. Annals of Surgical Oncology 2010 17 263–270.
(https://doi. org/10.1245/s10434-009-0716-x)

93 Postlewait LM, Ethun CG, Tran TB, Prescott JD, Pawlik TM, Wang TS, Glenn J,
Hatzaras I, Shenoy R, Phay JE et al. Outcomes of adjuvant mitotane after resection of
adrenocortical carcinoma: a 13-institution study by the US Adrenocortical Carcinoma Group.
Journal of the American College of Surgeons 2016 222 480–490. (https://doi.
org/10.1016/j.jamcollsurg.2015.12.013)

2356 94 Terzolo M, Angeli A, Fassnacht M, Daffara F, Tauchmanova L, Conton PA, Rossetto
2357 R, Buci L, Sperone P, Grossrubatscher E et al. Adjuvant mitotane treatment for
2358 adrenocortical carcinoma. New England Journal of Medicine 2007 356 2372–2380.
2359 (https://doi. org/10.1056/NEJMoa063360)

- 95 Bosco JL, Silliman RA, Thwin SS, Geiger AM, Buist DS, Prout MN, Yood MU, Haque
 R, Wei F & Lash TL. A most stubborn bias: no adjustment method fully resolves confounding
 by indication in observational studies. Journal of Clinical Epidemiology 2010 63 64–74.
 (https://doi.org/10.1016/j.jclinepi.2009.03.001)
- Hernan MA & Robins JM. Instruments for causal inference: an epidemiologist's
 dream? Epidemiology 2006 17 360–372. (https://doi.
 org/10.1097/01.ede.0000222409.00878.37)
- 2367 97 Suissa S. Immortal time bias in pharmaco-epidemiology. American Journal of 2368 Epidemiology 2008 167 492–499. (https://doi.org/10.1093/ aje/kwm324)
- 2369 98 Fassnacht M, Hahner S, Polat B, Koschker AC, Kenn W, Flentje M & Allolio B.
 2370 Efficacy of adjuvant radiotherapy of the tumor bed on local recurrence of adrenocortical
 2371 carcinoma. Journal of Clinical Endocrinology and Metabolism 2006 91 4501–4504.
- 99 Habra MA, Ejaz S, Feng L, Das P, Deniz F, Grubbs EG, Phan AT, Waguespack S,
 Montserrat AR, Jimenez C et al. A Retrospective Cohort Analysis of the Efficacy of Adjuvant
 Radiotherapy after Primary Surgical Resection in Patients with Adrenocortical Carcinoma.
 Journal of Clinical Endocrinology and Metabolism 2013 98 192–197.
 (https://doi.org/10.1210/jc.2012-2367)
- 2377 100 Sabolch A, Else T, Griffith KA, Ben-Josef E, Williams A, Miller BS, Worden F,
 2378 Hammer GD & Jolly S. Adjuvant radiation therapy improves local control after surgical
 2379 resection in patients with localized adrenocortical carcinoma. International Journal of
 2380 Radiation Oncology, Biology, Physics 2015 92 252–259. (https://doi.
 2381 org/10.1016/j.ijrobp.2015.01.007)
- 101 Berruti A, Terzolo M, Sperone P, Pia A, Della Casa S, Gross DJ, Carnaghi C, Casali
 P, Porpiglia F, Mantero F et al. Etoposide, doxorubicin and cisplatin plus mitotane in the
 treatment of advanced adrenocortical carcinoma: a large prospective phase II trial.
 Endocrine-Related Cancer 2005 12 657–666. (https://doi.org/10.1677/ erc.1.01025)
- 102 Fassnacht M, Berruti A, Baudin E, Demeure MJ, Gilbert J, Haak H, Kroiss M, Quinn
 DI, Hesseltine E, Ronchi CL et al. Linsitinib (OSI- 906) versus placebo for patients with
 locally advanced or metastatic adrenocortical carcinoma: a double-blind, randomised, phase
 3 study. Lancet Oncology 2015 16 426–435. (https://doi.org/10.1016/ S1470-2045(15)700811)
- 103 Hermsen IG, Fassnacht M, Terzolo M, Houterman S, den Hartigh J, Leboulleux S,
 Daffara F, Berruti A, Chadarevian R, Schlumberger M et al. Concentrations of o,p' DDD, o,p'
 DDA, and o,p' DDE as predictors of tumor response to mitotane in adrenocortical carcinoma:
 results of a retrospective ENS@T multicenter study. Journal of Clinical Endocrinology and
 Metabolism 2011 96 1844–1851.
- 104 Sperone P, Ferrero A, Daffara F, Priola A, Zaggia B, Volante M, Santini D, Vincenzi
 B, Badalamenti G, Intrivici C et al. Gemcitabine plus metronomic 5-fluorouracil or
 capecitabine as a second-/third-line chemotherapy in advanced adrenocortical carcinoma: a
 multicenter phase II study. Endocrine-Related Cancer 2010 17 445–453.
 (https://doi.org/10.1677/ERC-09-0281)
- Abraham J, Bakke S, Rutt A, Meadows B, Merino M, Alexander R, Schrump D,
 Bartlett D, Choyke P, Robey R et al. A phase II trial of combination chemotherapy and
 surgical resection for the treatment of metastatic adrenocortical carcinoma: continuous
 infusion doxorubicin, vincristine, and etoposide with daily mitotane as a P-glycoprotein
 antagonist. Cancer 2002 94 2333–2343. (https://doi. org/10.1002/cncr.10487)

- 2406 106 Baudin E, Docao C, Gicquel C, Vassal G, Bachelot A, Penfornis A & Schlumberger
 2407 M. Use of a topoisomerase I inhibitor (irinotecan, CPT-11) in metastatic adrenocortical
 2408 carcinoma. Annals of Oncology 2002 13 1806–1809.
- 2409 (https://doi.org/10.1093/annonc/mdf291)
- 2410 107 Baudin E, Pellegriti G, Bonnay M, Penfornis A, Laplanche A, Vassal G & Schlumberger
- M. Impact of monitoring plasma 1,1-dichlorodiphenildichloroethane (o,p' DDD) levels on the
- treatment of patients with adrenocortical carcinoma. Cancer 2001 92 1385–1392.
- 2413 (https://doi.org/10.1002/1097- 0142(20010915)92:6<1385::AID-CNCR1461>3.0.CO;2-2)
- 2414 108 Berruti A, Sperone P, Ferrero A, Germano A, Ardito A, Priola AM, De Francia S,
 2415 Volante M, Daffara F, Generali D et al. Phase II study of weekly paclitaxel and sorafenib as
 2416 second/third-line therapy in patients with adrenocortical carcinoma. European Journal of
 2417 Endocrinology 2012 166 451–458. (https://doi.org/10.1530/EJE-11-0918)
- 2418 109 Bonacci R, Gigliotti A, Baudin E, Wion-Barbot N, Emy P, Bonnay M, Cailleux AF,
 2419 Nakib I & Schlumberger M. Cytotoxic therapy with etoposide and cisplatin in advanced
 2420 adrenocortical carcinoma. British Journal of Cancer 1998 78 546–549.
 2421 (https://doi.org/10.1038/ bjc.1998.530)
- 2422 110 Bukowski RM, Wolfe M, Levine HS, Crawford DE, Stephens RL, Gaynor E & Harker
 2423 WG. Phase II trial of mitotane and cisplatin in patients with adrenal carcinoma: a Southwest
 2424 Oncology Group study. Journal of Clinical Oncology 1993 11 161–165. (https://doi.
 2425 org/10.1200/JCO.1993.11.1.161)
- 2426 111 Decker RA, Elson P, Hogan TF, Citrin DL, Westring DW, Banerjee TK, Gilchrist KW &
 2427 Horton J. Eastern-Cooperative-Oncology-Group Study 1879 mitotane and adriamycin in
 2428 patients with advanced adrenocortical carcinoma. Surgery 1991 110 1006–1013.
- Haak HR, Hermans J, van de Velde CJ, Lentjes EG, Goslings BM, Fleuren GJ &
 Krans HM. Optimal treatment of adrenocortical carcinoma with mitotane: results in a
 consecutive series of 96 patients. British Journal of Cancer 1994 69 947–951. (https://doi.
 org/10.1038/bjc.1994.183)
- Haluska P, Worden F, Olmos D, Yin D, Schteingart D, Batzel GN, Paccagnella ML,
 de Bono JS, Gualberto A & Hammer GD. Safety, tolerability, and pharmacokinetics of the
 anti-IGF-1R monoclonal antibody figitumumab in patients with refractory adrenocortical
 carcinoma. Cancer Chemotherapy and Pharmacology 2010 65 765–773.
 (https://doi.org/10.1007/s00280-009-1083-9)
- 2438 114 Khan TS, Sundin A, Juhlin C, Wilander E, Oberg K & Eriksson B. Vincristine,
 2439 cisplatin, teniposide, and cyclophosphamide combination in the treatment of recurrent or
 2440 metastatic adrenocortical cancer. Medical Oncology 2004 21 167–177. (https://
 2441 doi.org/10.1385/MO:21:2:167)
- Kroiss M, Deutschbein T, Schlotelburg W, Ronchi CL, Neu B, Muller HH, Quinkler M,
 Hahner S, Heidemeier A & Fassnacht M. Salvage treatment of adrenocortical carcinoma with
 trofosfamide. Hormones and Cancer 2016 7 211–218. (https://doi.org/10.1007/ s12672-0160260-7)
- Kroiss M, Quinkler M, Johanssen S, van Erp NP, Lankheet N, Pollinger A, Laubner K,
 Strasburger CJ, Hahner S, Muller HH et al. Sunitinib in refractory adrenocortical carcinoma: a
 phase II, single- arm, open-label trial. Journal of Clinical Endocrinology and Metabolism 2012
 97 3495–3503. (https://doi.org/10.1210/jc.2012-1419)

- 117 Naing A, LoRusso P, Fu S, Hong D, Chen HX, Doyle LA, Phan AT, Habra MA &
 2451 Kurzrock R. Insulin growth factor receptor (IGF-1R) antibody cixutumumab combined with
 2452 the mTOR inhibitor temsirolimus in patients with metastatic adrenocortical carcinoma. British
 2453 Journal of Cancer 2013 108 826–830. (https://doi.org/10.1038/ bjc.2013.46)
- 2454 118 O'Sullivan C, Edgerly M, Velarde M, Wilkerson J, Venkatesan AM, Pittaluga S, Yang
 2455 SX, Nguyen D, Balasubramaniam S & Fojo T. The VEGF inhibitor axitinib has limited
 2456 effectiveness as a therapy for adrenocortical cancer. Journal of Clinical Endocrinology and
 2457 Metabolism 2014 99 1291–1297.
- 2458 119 Quinkler M, Hahner S, Wortmann S, Johanssen S, Adam P, Ritter C, Strasburger C,
 2459 Allolio B & Fassnacht M. Treatment of advanced adrenocortical carcinoma with erlotinib plus
 2460 gemcitabine. Journal of Clinical Endocrinology and Metabolism 2008 93 2057–2062. (https://
 2461 doi.org/10.1210/jc.2007-2564)
- Schlumberger M, Brugieres L, Gicquel C, Travagli JP, Droz JP & Parmentier C. 5Fluorouracil, doxorubicin, and cisplatin as treatment for adrenal-cortical carcinoma. Cancer
 1991 67 2997–3000. (https://doi.org/10.1002/1097-0142(19910615)67:12<2997::AID-
 CNCR2820671211>3.0.CO;2-#)
- 2466 121 Urup T, Pawlak WZ, Petersen PM, Pappot H, Rorth M & Daugaard G. Treatment with
 2467 docetaxel and cisplatin in advanced adrenocortical carcinoma, a phase II study. British
 2468 Journal of Cancer 2013 108 1994–1997. (https://doi.org/10.1038/bjc.2013.229)
- Williamson SK, Lew D, Miller GJ, Balcerzak SP, Baker LH & Crawford ED. Phase II
 evaluation of cisplatin and etoposide followed by mitotane at disease progression in patients
 with locally advanced or metastatic adrenocortical carcinoma a Southwest Oncology Group
 study. Cancer 2000 88 1159–1165. (https://doi.org/10.1002/ (SICI)10970142(20000301)88:5<1159::AID-CNCR28>3.0.CO;2-R)
- 2474 123 Wortmann S, Quinkler M, Ritter C, Kroiss M, Johanssen S, Hahner S, Allolio B &
 2475 Fassnacht M. Bevacizumab plus capecitabine as a salvage therapy in advanced
 2476 adrenocortical carcinoma. European Journal of Endocrinology 2010 162 349–356.
 2477 (https://doi.org/10.1530/EJE-09-0804)
- Henning JEK, Deutschbein T, Altieri B, Steinhauer S, Kircher S, Sbiera S, Wild V,
 Schlotelburg W, Kroiss M, Perotti P et al. Gemcitabine-based chemotherapy in
 adrenocortical carcinoma: a multicenter study of efficacy and predictive factors. Journal of
 Clinical Endocrinology and Metabolism 2017 102 4323–4332. (https://doi.
 org/10.1210/jc.2017-01624)
- Lerario AM, Worden FP, Ramm CA, Hesseltine EA, Stadler WM, Else T, Shah MH,
 Agamah E, Rao K & Hammer GD. The combination of insulin-like growth factor receptor 1
 (IGF1R) antibody cixutumumab and mitotane as a first-line therapy for patients with
 recurrent/metastatic adrenocortical carcinoma: a multi-institutional NCI-sponsored trial.
 Hormones and Cancer 2014 5 232–239. (https:// doi.org/10.1007/s12672-014-0182-1)
- 2488 126 Datrice NM, Langan RC, Ripley RT, Kemp CD, Steinberg SM, Wood BJ, Libutti SK,
 2489 Fojo T, Schrump DS & Avital I. Operative management for recurrent and metastatic
 2490 adrenocortical carcinoma. Journal of Surgical Oncology 2012 105 709–713. (https://doi.
 2491 org/10.1002/jso.23015)
- 2492 127 Gaujoux S, Al-Ahmadie H, Allen PJ, Gonen M, Shia J, D'Angelica M, Dematteo R,
 2493 Fong Y, Blumgart L & Jarnagin WR. Resection of adrenocortical carcinoma liver metastasis:
 2494 is it justified? Annals of Surgical Oncology 2012 19 2643–2651. (https://doi.org/10.1245/
 2495 s10434-012-2358-7)

2496 128 Kemp CD, Ripley RT, Mathur A, Steinberg SM, Nguyen DM, Fojo T & Schrump DS.
2497 Pulmonary resection for metastatic adrenocortical carcinoma: the National Cancer Institute
2498 experience. Annals of Thoracic Surgery 2011 92 1195–1200. (https://doi.org/10.1016/j.
2499 athoracsur.2011.05.013)

129 Kwauk S & Burt M. Pulmonary metastases from adrenal cortical carcinoma: results of
resection. Journal of Surgical Oncology 1993 53 243–246.
(https://doi.org/10.1002/jso.2930530411)

2503 130 op den Winkel J, Pfannschmidt J, Muley T, Grunewald C, Dienemann H, Fassnacht M
2504 & Allolio B. Metastatic adrenocortical carcinoma: results of 56 pulmonary metastasectomies
2505 in 24 patients. Annals of Thoracic Surgery 2011 92 1965–1970. (https://doi.
2506 org/10.1016/j.athoracsur.2011.07.088)

131 Ripley RT, Kemp CD, Davis JL, Langan RC, Royal RE, Libutti SK, Steinberg SM,
Wood BJ, Kammula US, Fojo T et al. Liver resection and ablation for metastatic
adrenocortical carcinoma. Annals of Surgical Oncology 2011 18 1972–1979.
(https://doi.org/10.1245/ s10434-011-1564-z)

Bellantone R, Ferrante A, Boscherini M, Lombardi CP, Crucitti P, Crucitti F, Favia G,
Borrelli D, Boffi L, Capussotti L et al. Role of reoperation in recurrence of adrenal cortical
carcinoma: results from 188 cases collected in the Italian National Registry for Adrenal
Cortical Carcinoma. Surgery 1997 122 1212–1218. (https://doi. org/10.1016/S0039-

2515 6060(97)90229-4)

2516 133 Crucitti F, Bellantone R, Ferrante A, Boscherini M, Crucitti P, Carbone G, Casaccia
2517 M, Campisi C, Cavallaro A, Sapienza P et al. The Italian registry for adrenal cortical
2518 carcinoma: analysis of a multiinstitutional series of 129 patients. Surgery 1996 119 161–170.
2519 (https://doi.org/10.1016/S0039-6060(96)80164-4)

134 Dy BM, Wise KB, Richards ML, Young WE, Grant CS, Bible KC, Rosedahl J,
Harmsen WS, Farley DR & Thompson GB. Operative intervention for recurrent
adrenocortical cancer. Surgery 2013 154 1292–1299.
(https://doi.org/10.1016/i.surg.2013.06.033)

2523 (https://doi.org/10.1016/j.surg.2013.06.033)

- 135 Jensen JC, Pass HI, Sindelar WF & Norton JA. Recurrent or metastatic disease in
 select patients with adrenocortical carcinoma aggressive resection vs chemotherapy.
 Archives of Surgery 1991 126 457–461.
- 2527 (https://doi.org/10.1001/archsurg.1991.01410280059008)
- Simon G, Pattou F, Mirallie E, Lifante JC, Nomine C, Arnault V, de Calan L, Gaillard
 C, Carnaille B, Brunaud L et al. Surgery for recurrent adrenocortical carcinoma: a multicenter
 retrospective study. Surgery 2017 161 249–255. (https://doi.org/10.1016/j. surg.2016.08.058)
- 137 Tran TB, Liou D, Menon VG & Nissen NN. Surgical management of advanced
 adrenocortical carcinoma: a 21-year population-based analysis. American Surgeon 2013 79
 1115–1118.
- 2534 138 Dy BM, Strajina V, Cayo AK, Richards ML, Farley DR, Grant CS, Harmsen WS,
- 2535 Evans DB, Grubbs EG, Bible KC et al. Surgical resection of synchronously metastatic
- adrenocortical cancer. Annals of Surgical Oncology 2015 22 146–151.
- 2537 (https://doi.org/10.1245/s10434-014-3944-7)
- 139 Hahner S, Kreissl MC, Fassnacht M, Haenscheid H, Knoedler P, Lang K, Buck AK,
 Reiners C, Allolio B & Schirbel A. [131] Iodometomidate for targeted radionuclide therapy of

- advanced adrenocortical carcinoma. Journal of Clinical Endocrinology and Metabolism 2012
 97 914–922. (https://doi.org/10.1210/jc.2011-2765)
- 140 Cazejust J, De Baere T, Auperin A, Deschamps F, Hechelhammer L, Abdel-Rehim M,
 Schlumberger M, Leboulleux S & Baudin E. Transcatheter arterial chemoembolization for
 liver metastases in patients with adrenocortical carcinoma. Journal of Vascular and
 Interventional Radiology 2010 21 1527–1532.
- 141 Wood BJ, Abraham J, Hvizda JL, Alexander HR & Fojo T. Radiofrequency ablation of
 adrenal tumors and adrenocortical carcinoma metastases. Cancer 2003 97 554–560.
 (https://doi. org/10.1002/cncr.11084)
- Ho J, Turkbey B, Edgerly M, Alimchandani M, Quezado M, Camphausen K, Fojo T &
 Kaushal A. Role of radiotherapy in adrenocortical carcinoma. Cancer Journal 2013 19 288–
 294. (https://doi.org/10.1097/PPO.0b013e31829e3221)
- 2552 143 Hahner S & Fassnacht M. Mitotane for adrenocortical carcinoma treatment. Current 2553 Opinion in Investigational Drugs 2005 6 386–394.
- 2554144Stell A & Sinnott R. The ENSAT registry: a digital repository supporting adrenal2555cancer research. Studies in Health Technology and Informatics 2012 178 207–212.
- 145 Fassnacht M, Kenn W & Allolio B. Adrenal tumors: how to establish malignancy?
 Journal of Endocrinological Investigation 2004 27 387–399.
 (https://doi.org/10.1007/BF03351068)
- 146 Allolio B & Fassnacht M. Clinical review: adrenocortical carcinoma: clinical update.
 Journal of Clinical Endocrinology and Metabolism 2006 91 2027–2037.
 (https://doi.org/10.1210/jc.2005-2639)
- 147 Nieman LK, Biller BM, Findling JW, Newell-Price J, Savage MO, Stewart PM &
 2563 Montori VM. The diagnosis of Cushing's syndrome: an Endocrine Society Clinical Practice
 2564 Guideline. Journal of Clinical Endocrinology and Metabolism 2008 93 1526–1540.
 2565 (https://doi. org/10.1210/jc.2008-0125)
- Libe R, Fratticci A & Bertherat J. Adrenocortical cancer: pathophysiology and clinical
 management. Endocrine-Related Cancer 2007 14 13–28.
 (https://doi.org/10.1677/erc.1.01130)
- Legro RS, Arslanian SA, Ehrmann DA, Hoeger KM, Murad MH, Pasquali R & Welt
 CK. Diagnosis and treatment of polycystic ovary syndrome: an Endocrine Society clinical
 practice guideline. Journal of Clinical Endocrinology and Metabolism 2013 98 4565–4592.
 (https:// doi.org/10.1210/jc.2013-2350)
- 150 Funder JW, Carey RM, Mantero F, Murad MH, Reincke M, Shibata H, Stowasser M &
 Young WF Jr. The management of primary aldosteronism: case detection, diagnosis, and
 treatment: an Endocrine Society Clinical Practice Guideline. Journal of Clinical Endocrinology
 and Metabolism 2016 101 1889–1916. (https://doi. org/10.1210/jc.2015-4061)
- 2577 151 Dinnes J, Bancos I, Ferrante di Ruffano L, Chortis V, Davenport C, Bayliss S, Sahdev
 2578 A, Guest P, Fassnacht M, Deeks JJ et al. MANAGEMENT OF ENDOCRINE DISEASE:
 2579 Imaging for the diagnosis of malignancy in incidentally discovered adrenal masses: a
 2580 systematic review and meta-analysis. European Journal of Endocrinology 2016 175 R51–
 2581 R64. (https://doi.org/10.1530/EJE-16-0461)

- 2582 152 Peppercorn PD, Grossman AB & Reznek RH. Imaging of incidentally discovered
 2583 adrenal masses. Clinical Endocrinology 1998 48 379–388. (https://doi.org/10.1046/j.13652265.1998.00475.x)
- 2585 153 Caoili EM, Korobkin M, Francis IR, Cohan RH, Platt JF, Dunnick NR & Raghupathi KI.
 2586 Adrenal masses: characterization with combined unenhanced and delayed enhanced CT.
 2587 Radiology 2002 222 629–633. (https://doi.org/10.1148/radiol.2223010766)
- Blake MA, Kalra MK, Sweeney AT, Lucey BC, Maher MM, Sahani DV, Halpern EF,
 Mueller PR, Hahn PF & Boland GW. Distinguishing benign from malignant adrenal masses:
 multi-detector row CT protocol with 10-minute delay. Radiology 2006 238 578–585. (https://
 doi.org/10.1148/radiol.2382041514)
- 155 Ilias I, Sahdev A, Reznek RH, Grossman AB & Pacak K. The optimal imaging of
 adrenal tumours: a comparison of different methods. Endocrine-Related Cancer 2007 14
 587–599. (https://doi.org/10.1677/ ERC-07-0045)
- 156 Mackie GC, Shulkin BL, Ribeiro RC, Worden FP, Gauger PG, Mody RJ, Connolly LP,
 Kunter G, Rodriguez-Galindo C, Wallis JW et al. Use of [18F]fluorodeoxyglucose positron
 emission tomography in evaluating locally recurrent and metastatic adrenocortical
 carcinoma. Journal of Clinical Endocrinology and Metabolism 2006 91 2665–2671.
 (https://doi.org/10.1210/jc.2005-2612)
- 2600 157 Groussin L, Bonardel G, Silvera S, Tissier F, Coste J, Abiven G, Libe R, Bienvenu M,
 2601 Alberini JL, Salenave S et al. 18F-Fluorodeoxyglucose positron emission tomography for the
 2602 diagnosis of adrenocortical tumors: a prospective study in 77 operated patients. Journal of
 2603 Clinical Endocrinology and Metabolism 2009 94 1713–1722. (https://
 2604 doi.org/10.1210/jc.2008-2302)
- 2605 158 Deandreis D, Leboulleux S, Caramella C, Schlumberger M & Baudin E. FDG PET in
 2606 the management of patients with adrenal masses and adrenocortical carcinoma. Hormones
 2607 and Cancer 2011 2 354–362. (https://doi.org/10.1007/s12672-011-0091-5)
- 2608 159 Cistaro A, Niccoli Asabella A, Coppolino P, Quartuccio N, Altini C, Cucinotta M,
 2609 Alongi P, Balma M, Sanfilippo S, Buschiazzo A et al. Diagnostic and prognostic value of 18F2610 FDG PET/CT in comparison with morphological imaging in primary adrenal gland
 2611 malignancies a multicenter experience. Hellenic Journal of Nuclear Medicine 2015 18 97–
 2612 102.
- Altinmakas E, Hobbs BP, Ye H, Grubbs EG, Perrier ND, Prieto VG, Lee JE & Ng CS.
 Diagnostic performance of (18-)F-FDG-PET-CT in adrenal lesions using histopathology as
 reference standard. Abdominal Radiology 2017 42 577–584. (https://doi.org/10.1007/
 s00261-016-0915-4)
- 2617 161 Ciftci E, Turgut B, Cakmakcilar A & Erturk SA. Diagnostic importance of 18F-FDG
 2618 PET/CT parameters and total lesion glycolysis in differentiating between benign and
 2619 malignant adrenal lesions. Nuclear Medicine Communications 2017 38 788–794. (https://doi.
 2620 org/10.1097/MNM.00000000000712)
- 162 Bluemel C, Hahner S, Heinze B, Fassnacht M, Kroiss M, Bley TA, Wester HJ, Kropf
 S, Lapa C, Schirbel A et al. Investigating the chemokine receptor 4 as potential theranostic
 target in adrenocortical cancer patients. Clinical Nuclear Medicine 2017 42 e29–e34.
 (https://doi.org/10.1097/RLU.00000000001435)
- 2625 163 Werner RA, Kroiss M, Nakajo M, Mugge DO, Hahner S, Fassnacht M, Schirbel A,
 2626 Bluemel C, Higuchi T, Papp L et al. Assessment of tumor heterogeneity in treatment-naive

adrenocortical cancer patients using 18F-FDG positron emission tomography. Endocrine
2016 53 791–800. (https://doi.org/10.1007/s12020-016-0970-1)

164 Wu YW & Tan CH. Determination of a cutoff attenuation value on single-phase
contrast-enhanced CT for characterizing adrenal nodules via chemical shift MRI. Abdominal
Radiology 2016 41 1170–1177. (https://doi.org/10.1007/s00261-016-0654-6)

165 Nakajo M, Jinguji M, Shinaji T, Nakabeppu Y, Fukukura Y & Yoshiura T. Texture
analysis of FDG PET/CT for differentiating between FDG-avid benign and metastatic adrenal
tumors: efficacy of combining SUV and texture parameters. Abdominal Radiology 2017 42
2882–2889. (https://doi.org/10.1007/s00261-017-1207-3)

- 166 Guerin C, Pattou F, Brunaud L, Lifante JC, Mirallie E, Haissaguerre M, Huglo D,
 Olivier P, Houzard C, Ansquer C et al. Performance of 18F- FDG PET/CT in the
 characterization of adrenal masses in noncancer patients: a prospective study. Journal of
 Clinical Endocrinology and Metabolism 2017 102 2465–2472.
 (https://doi.org/10.1210/jc.2017-00254)
- 167 Marty M, Gaye D, Perez P, Auder C, Nunes ML, Ferriere A, Haissaguerre M &
 Tabarin A. Diagnostic accuracy of computed tomography to identify adenomas among
 adrenal incidentalomas in an endocrinological population. European Journal of
 Endocrinology 2018 178 439–446. (https://doi.org/10.1530/EJE-17-1056)
- 168 Kim SJ, Lee SW, Pak K, Kim IJ & Kim K. Diagnostic accuracy of (18) F-FDG PET or
 2646 PET/CT for the characterization of adrenal masses: a systematic review and meta-analysis.
 2647 British Journal of Radiology 2018 91 20170520. (https://doi.org/10.1259/bjr.20170520)
- 169 Delivanis DA, Bancos I, Atwell TD, Schmit GD, Eiken PW, Natt N, Erickson D,
 2649 Maraka S, Young WF & Nathan MA. Diagnostic performance of unenhanced computed
 2650 tomography and (18) F-fluorodeoxyglucose positron emission tomography in indeterminate
 2651 adrenal tumours. Clinical Endocrinology 2018 88 30–36. (https://doi.org/10.1111/cen.13448)
- 2652 170 Romeo V, Maurea S, Cuocolo R, Petretta M, Mainenti PP, Verde F, Coppola M,
 2653 Dell'Aversana S & Brunetti A. Characterization of adrenal lesions on unenhanced MRI using
 2654 texture analysis: a machine-learning approach. Journal of Magnetic Resonance Imaging
 2655 2018 48 198–204. (https://doi.org/10.1002/jmri.25954)
- 171 Thomas AJ, Habra MA, Bhosale PR, Qayyum AA, Ahmed K, Vicens R & Elsayes KM.
 1057 Interobserver agreement in distinguishing large adrenal adenomas and adrenocortical
 carcinomas on computed tomography. Abdominal Radiology 2018 Epub. (https://doi.
 org/10.1007/s00261-018-1603-3)
- 172 Ng CS, Altinmakas E, Wei W, Ghosh P, Li X, Grubbs EG, Perrier NA, Prieto VG, Lee
 JE & Hobbs BP. Combining washout and noncontrast data from adrenal protocol CT:
 improving diagnostic performance. Academic Radiology 2018 25 861–868.
 (https://doi.org/10.1016/j. acra.2017.12.005)
- Petersenn S, Richter PA, Broemel T, Ritter CO, Deutschbein T, Beil FU, Allolio B &
 Fassnacht M. Computed tomography criteria for discrimination of adrenal adenomas and
 adrenocortical carcinomas: analysis of the German ACC registry. European Journal of
 Endocrinology 2015 172 415–422. (https://doi.org/10.1530/EJE-14-0916)
- 174 Bancos I, Tamhane S, Shah M, Delivanis DA, Alahdab F, Arlt W, Fassnacht M &
 2669 Murad MH. DIAGNOSIS OF ENDOCRINE DISEASE: The diagnostic performance of adrenal
 2670 biopsy: a systematic review and meta-analysis. European Journal of Endocrinology 2016 175
 2671 R65–R80. (https://doi.org/10.1530/EJE-16-0297)

- 2672 175 Williams AR, Hammer GD & Else T. Transcutaneous biopsy of adrenocortical
 2673 carcinoma is rarely helpful in diagnosis, potentially harmful, but does not affect patient
 2674 outcome. European Journal of Endocrinology 2014 170 829–835.
 2675 (https://doi.org/10.1520/5.15.40.4000)
- 2675 (https://doi.org/10.1530/EJE-13-1033)

2676 176 Palazzo F, Dickinson A, Phillips B, Sahdev A, Bliss R, Rasheed A, Krukowski Z &
2677 Newell-Price J. Adrenal surgery in England: better outcomes in high-volume practices.
2678 Clinical Endocrinology 2016 85 17–20. (https://doi.org/10.1111/cen.13021)

- 2679 177 Park HS, Roman SA & Sosa JA. Outcomes from 3144 adrenalectomies in the United
 2680 States: which matters more, surgeon volume or specialty? Archives of Surgery 2009 144
 2681 1060–1067. (https://doi. org/10.1001/archsurg.2009.191)
- 2682 178 Lindeman B, Hashimoto DA, Bababekov YJ, Stapleton SM, Chang DC, Hodin RA &
 2683 Phitayakorn R. Fifteen years of adrenalectomies: impact of specialty training and operative
 2684 volume. Surgery 2018 163 150–156. (https://doi.org/10.1016/j. surg.2017.05.024)
- Villar JM, Moreno P, Ortega J, Bollo E, Ramirez CP, Munoz N, Martinez C,
 Dominguez-Adame E, Sancho J, Del Pino JM et al. Results of adrenal surgery. Data of a
 Spanish National Survey. Langenbecks Archives of Surgery 2010 395 837–843.
 (https://doi.org/10.1007/ s00423-010-0697-z)
- 180 Gallagher SF, Wahi M, Haines KL, Baksh K, Enriquez J, Lee TM, Murr MM & Fabri
 PJ. Trends in adrenalectomy rates, indications, and physician volume: a statewide analysis
 of 1816 adrenalectomies. Surgery 2007 142 1011–1021; discussion 1011–1021. (https://doi.
 org/10.1016/j.surg.2007.09.024)
- 2693 181 Lombardi CP, Raffaelli M, Boniardi M, De Toma G, Marzano LA, Miccoli P, Minni F,
 2694 Morino M, Pelizzo MR, Pietrabissa A et al. Adrenocortical carcinoma: effect of hospital
 2695 volume on patient outcome. Langenbecks Archives of Surgery 2012 397 201–207. (https://
 2696 doi.org/10.1007/s00423-011-0866-8)
- 2697 182 Gratian L, Pura J, Dinan M, Reed S, Scheri R, Roman S & Sosa JA. Treatment
 2698 patterns and outcomes for patients with adrenocortical carcinoma associated with hospital
 2699 case volume in the United States. Annals of Surgical Oncology 2014 21 3509–3514.
 2700 (https://doi. org/10.1245/s10434-014-3931-z)
- 183 Hermsen IG, Kerkhofs TM, den Butter G, Kievit J, van Eijck CH, Nieveen van Dijkum
 EJ & Haak HR. Surgery in adrenocortical carcinoma: Importance of national cooperation and
 centralized surgery. Surgery 2012 152 50–56. (https://doi.org/10.1016/j. surg.2012.02.005)
- 184 Kerkhofs TM, Verhoeven RH, Bonjer HJ, van Dijkum EJ, Vriens MR, De Vries J, Van
 Eijck CH, Bonsing BA, Van de Poll-Franse LV & Haak HR. Surgery for adrenocortical
 carcinoma in The Netherlands: analysis of the national cancer registry data. European
 Journal of Endocrinology 2013 169 83–89. (https://doi.org/10.1530/EJE-13-0142)
- 185 Gaujoux S & Mihai R. European Society of Endocrine Surgeons (ESES) and
 European Network for the Study of Adrenal Tumours (ENSAT) recommendations for the
 surgical management of adrenocortical carcinoma. British Journal of Surgery 2017 104 358–
 376. (https://doi. org/10.1002/bjs.10414)
- 186 Gaujoux S & Brennan MF. Recommendation for standardized surgical management
 of primary adrenocortical carcinoma. Surgery 2012 152 123–132.
- 2714 (https://doi.org/10.1016/j.surg.2011.09.030)
- 2715 187 Porpiglia F, Fiori C, Daffara FC, Zaggia B, Ardito A, Scarpa RM, Papotti M, Berruti A,
 2716 Scagliotti GV & Terzolo M. Does nephrectomy during radical adrenalectomy for stage II

- adrenocortical cancer affect patient outcome? Journal of Endocrinological Investigation 2016
 39 465–471. (https://doi.org/10.1007/s40618-015-0422-4)
- 188 Donatini G, Caiazzo R, Do Cao C, Aubert S, Zerrweck C, El-Kathib Z, Gauthier T,
 Leteurtre E, Wemeau JL, Vantyghem MC et al. Long-term survival after adrenalectomy for
 stage I/II adrenocortical carcinoma (ACC): a retrospective comparative cohort study of
 laparoscopic versus open approach. Annals of Surgical Oncology 2014 21 284–291.
 (https://doi.org/10.1245/s10434-013-3164-6)
- 189 Sgourakis G, Lanitis S, Kouloura A, Zaphiriadou P, Karkoulias K, Raptis D,
 Anagnostara A & Caraliotas C. Laparoscopic versus open adrenalectomy for stage I/II
 adrenocortical carcinoma: meta-analysis of outcomes. Journal of Investigative Surgery 2015
 28 145–152. (https://doi.org/10.3109/08941939.2014.987886)
- Autorino R, Bove P, De Sio M, Miano R, Micali S, Cindolo L, Greco F, Nicholas J,
 Fiori C, Bianchi G et al. Open versus laparoscopic adrenalectomy for adrenocortical
 carcinoma: a meta-analysis of surgical and oncological outcomes. Annals of Surgical
 Oncology 2016 23 1195–1202. (https://doi.org/10.1245/s10434-015-4900-x)
- 2732 191 Langenhuijsen J, Birtle A, Klatte T, Porpiglia F & Timsit MO. Surgical management of
 adrenocortical carcinoma: impact of laparoscopic approach, lymphadenectomy, and surgical
 volume on outcomes-a systematic review and meta-analysis of the current literature.
 European Urology Focus 2016 1 241–250. (https://doi.org/10.1016/j. euf.2015.12.001)
- Lee CW, Salem AI, Schneider DF, Leverson GE, Tran TB, Poultsides GA, Postlewait
 LM, Maithel SK, Wang TS, Hatzaras I et al. Minimally invasive resection of adrenocortical
 carcinoma: a multi- institutional study of 201 patients. Journal of Gastrointestinal Surgery
 2017 21 352–362. (https://doi.org/10.1007/s11605-016-3262-4)
- 2740 193 Zheng GY, Li HZ, Deng JH, Zhang XB & Wu XC. Open adrenalectomy versus
 2741 laparoscopic adrenalectomy for adrenocortical carcinoma: a retrospective comparative study
 2742 on short-term oncologic prognosis. OncoTargets and Therapy 2018 11 1625–1632.
 2743 (https://doi.org/10.2147/OTT.S157518)
- 194 Mpaili E, Moris D, Tsilimigras DI, Oikonomou D, Pawlik TM, Schizas D, Papalampros
 A, Felekouras E & Dimitroulis D. Laparoscopic versus open adrenalectomy for
 localized/locally advanced primary adrenocortical carcinoma (ENSAT I-III) in adults: is
 margin-free resection the key surgical factor that dictates outcome? A review of the literature.
 Journal of Laparoendoscopic and Advanced Surgical Techniques Part A 2018 28 408–414.
 (https://doi.org/10.1089/ lap.2017.0546)
- Huynh KT, Lee DY, Lau BJ, Flaherty DC, Lee J & Goldfarb M. Impact of laparoscopic
 adrenalectomy on overall survival in patients with nonmetastatic adrenocortical carcinoma.
 Journal of the American College of Surgeons 2016 223 485–492. (https://doi.org/10.1016/j.
 jamcollsurg.2016.05.015)
- 196 Barczynski M, Konturek A, Golkowski F, Cichon S, Huszno B, Peitgen K & Walz MK.
 Posterior retroperitoneoscopic adrenalectomy: a comparison between the initial experience
 in the invention phase and introductory phase of the new surgical technique. World Journal
 of Surgery 2007 31 65–71. (https://doi.org/10.1007/s00268-006-0083-8)
- 2758 197 Schreinemakers JM, Kiela GJ, Valk GD, Vriens MR & Rinkes IH. Retroperitoneal
 2759 endoscopic adrenalectomy is safe and effective. British Journal of Surgery 2010 97 1667–
 2760 1672. (https://doi. org/10.1002/bjs.7191)

- 2761 198 Nilubol N, Patel D & Kebebew E. Does lymphadenectomy improve survival in patients
 2762 with adrenocortical carcinoma? A population- based study. World Journal of Surgery 2016 40
 2763 697–705. (https://doi. org/10.1007/s00268-015-3283-2)
- 2764 199 Saade N, Sadler C & Goldfarb M. Impact of regional lymph node dissection on
 2765 disease specific survival in adrenal cortical carcinoma. Hormone and Metabolic Research
 2766 2015 47 820–825. (https://doi. org/10.1055/s-0035-1549877)
- 2767 200 Harrison LE, Gaudin PB & Brennan MF. Pathologic features of prognostic
 2768 significance for adrenocortical carcinoma after curative resection. Archives of Surgery 1999
 2769 134 181–185. (https://doi. org/10.1001/archsurg.134.2.181)
- 201 Panjwani S, Moore MD, Gray KD, Finnerty BM, Beninato T, Brunaud L, Fahey TJ 3rd
 2771 & Zarnegar R. The impact of nodal dissection on staging in adrenocortical carcinoma. Annals
 2772 of Surgical Oncology 2017 24 3617–3623. (https://doi.org/10.1245/s10434-017-6064-3)
- 202 Gerry JM, Tran TB, Postlewait LM, Maithel SK, Prescott JD, Wang TS, Glenn JA,
 2774 Phay JE, Keplinger K, Fields RC et al. Lymphadenectomy for adrenocortical carcinoma: is
 2775 there a therapeutic benefit? Annals of Surgical Oncology 2016 23 708–713.
 2776 (https://doi.org/10.1245/s10434-016-5536-1)
- 2777 203 Reibetanz J, Jurowich C, Erdogan I, Nies C, Rayes N, Dralle H, Behrend M, Allolio B
 2778 & Fassnacht M. Impact of lymphadenectomy on the oncologic outcome of patients with
 2779 adrenocortical carcinoma. Annals of Surgery 2012 255 363–369. (https://doi.org/10.1097/
 2780 SLA.0b013e3182367ac3)
- 2781 204 Chiche L, Dousset B, Kieffer E & Chapuis Y. Adrenocortical carcinoma extending into
 2782 the inferior vena cava: presentation of a 15-patient series and review of the literature.
 2783 Surgery 2006 139 15–27. (https://doi.org/10.1016/j.surg.2005.05.014)
- 2784 205 Turbendian HK, Strong VE, Hsu M, Ghossein RA & Fahey TJ. Adrenocortical
 2785 carcinoma: the influence of large vessel extension. Surgery 2010 148 1057–1064.
 2786 (https://doi.org/10.1016/j. surg.2010.09.024)
- 2787 206 Mihai R, Iacobone M, Makay O, Moreno P, Frilling A, Kraimps JL, Soriano A, Villar del
 2788 Moral J, Barczynski M, Duran MC et al. Outcome of operation in patients with adrenocortical
 2789 cancer invading the inferior vena cava a European Society of Endocrine Surgeons (ESES)
 2790 survey. Langenbecks Archives of Surgery 2012 397 225–231. (https://
 2791 doi.org/10.1007/s00423-011-0876-6)
- 2792 207 Eller-Vainicher C, Morelli V, Salcuni AS, Battista C, Torlontano M, Coletti F, Iorio L,
 2793 Cairoli E, Beck-Peccoz P, Arosio M et al. Accuracy of several parameters of hypothalamic2794 pituitary-adrenal axis activity in predicting before surgery the metabolic effects of the removal
 2795 of an adrenal incidentaloma. European Journal of Endocrinology 2010 163 925–935.
 2796 (https://doi.org/10.1530/EJE-10-0602)
- 2797 208 Bornstein SR, Allolio B, Arlt W, Barthel A, Don-Wauchope A, Hammer GD, Husebye
 2798 ES, Merke DP, Murad MH, Stratakis CA et al. Diagnosis and treatment of primary adrenal
 2799 insufficiency: an Endocrine Society Clinical Practice Guideline. Journal of Clinical
 2800 Endocrinology and Metabolism 2016 101 364–389. (https://doi. org/10.1210/jc.2015-1710)
- 209 Duregon E, Volante M, Bollito E, Goia M, Buttigliero C, Zaggia B, Berruti A, Scagliotti
 2802 GV & Papotti M. Pitfalls in the diagnosis of adrenocortical tumors: a lesson from 300
 2803 consultation cases. Human Pathology 2015 46 1799–1807. (https://doi.org/10.1016/j.
 2804 humpath.2015.08.012)

- 2805 210 Sangoi AR, Fujiwara M, West RB, Montgomery KD, Bonventre JV, Higgins JP, Rouse
 2806 RV, Gokden N & McKenney JK. Immunohistochemical distinction of primary adrenal cortical
 2807 lesions from metastatic clear cell renal cell carcinoma: a study of 248 cases. American
 2808 Journal of Surgical Pathology 2011 35 678–686. (https://doi.
- 2809 org/10.1097/PAS.0b013e3182152629)
- 2810 211 Weissferdt A, Phan A, Suster S & Moran CA. Adrenocortical carcinoma: a
 2811 comprehensive immunohistochemical study of 40 cases. Applied Immunohistochemistry and
 2812 Molecular Morphology 2014 22 24–30. (https://doi.org/10.1097/PAI.0b013e31828a96cf)
- 2813 212 Tissier F, Aubert S, Leteurtre E, Alghuzlan A, Patey M, Decaussin M, Dousset L,
- 2814 Gobet F, Hoang C, Mazerolles C et al. Adrenocortical tumors (ACT): evaluation and
 2815 harmonization of the reading of the Weiss system criteria at the French level. Laboratory
 2816 Investigation 2010 90 133A.
- 2817 213 Tissier F, Aubert S, Leteurtre E, Al Ghuzlan A, Patey M, Decaussin M, Doucet L,
 2818 Gobet F, Hoang C, Mazerolles C et al. Adrenocortical tumors: improving the practice of the
 2819 weiss system through virtual microscopy a national program of the French Network INCa2820 COMETE. American Journal of Surgical Pathology 2012 36 1194–1201.
- 2821 (https://doi.org/10.1097/PAS.0b013e31825a6308)
- 2822 214 Weiss LM. Comparative histologic study of 43 metastasizing and nonmetastasizing
 2823 adrenocortical tumors. American Journal of Surgical Pathology 1984 8 163–169.
 2824 (https://doi.org/10.1097/00000478-198403000-00001)
- 2825 215 Weiss LM, Medeiros LJ & Vickery AL Jr. Pathologic features of prognostic
 2826 significance in adrenocortical carcinoma. American Journal of Surgical Pathology 1989 13
 2827 202–206. (https://doi. org/10.1097/00000478-198903000-00004)
- 2828 216 van Slooten H, Schaberg A, Smeenk D & Moolenaar AJ. Morphologic characteristics
 2829 of benign and malignant adrenocortical tumors. Cancer 1985 55 766–773.
 2830 (https://doi.org/10.1002/1097- 0142(19850215)55:4<766::AID-CNCR2820550414>3.0.CO;22831 7)
- 2832 217 Duregon E, Fassina A, Volante M, Nesi G, Santi R, Gatti G, Cappellesso R, Dalino
 2833 Ciaramella P, Ventura L, Gambacorta M et al. The reticulin algorithm for adrenocortical tumor
 2834 diagnosis: a multicentric validation study on 245 unpublished cases. American Journal of
 2835 Surgical Pathology 2013 37 1433–1440. (https://doi.org/10.1097/ PAS.0b013e31828d387b)
- 2836 218 Bisceglia M, Ludovico O, Di Mattia A, Ben-Dor D, Sandbank J, Pasquinelli G, Lau SK
 2837 & Weiss LM. Adrenocortical oncocytic tumors: report of 10 cases and review of the literature.
 2838 International Journal of Surgical Pathology 2004 12 231–243. (https://doi.
 2839 org/10.1177/106689690401200304)
- 2840 219 Duregon E, Volante M, Cappia S, Cuccurullo A, Bisceglia M, Wong DD, Spagnolo
 2841 DV, Szpak-Ulczok S, Bollito E, Daffara F et al. Oncocytic adrenocortical tumors: diagnostic
 2842 algorithm and mitochondrial DNA profile in 27 cases. American Journal of Surgical Pathology
 2843 2011 35 1882–1893. (https://doi.org/10.1097/ PAS.0b013e31822da401)
- Wong DD, Spagnolo DV, Bisceglia M, Havlat M, McCallum D & Platten MA.
 Oncocytic adrenocortical neoplasms a clinicopathologic study of 13 new cases
 emphasizing the importance of their recognition. Human Pathology 2011 42 489–499.
 (https://doi. org/10.1016/j.humpath.2010.08.010)

2848 221 Fuhrman SA, Lasky LC & Limas C. Prognostic significance of morphologic
2849 parameters in renal cell carcinoma. American Journal of Surgical Pathology 1982 6 655–663.
2850 (https://doi. org/10.1097/00000478-198210000-00007)

2851 222 Lu H, Papathomas TG, van Zessen D, Palli I, de Krijger RR, van der Spek PJ,
2852 Dinjens WN & Stubbs AP. Automated Selection of Hotspots (ASH): enhanced automated
2853 segmentation and adaptive step finding for Ki67 hotspot detection in adrenal cortical cancer.
2854 Diagnostic Pathology 2014 9 216. (https://doi.org/10.1186/s13000-014-0216-6)

- 2855 223 Papathomas TG, Pucci E, Giordano TJ, Lu H, Duregon E, Volante M, Papotti M,
 2856 Lloyd RV, Tischler AS, van Nederveen FH et al. An international Ki67 reproducibility study in
 2857 adrenal cortical carcinoma. American Journal of Surgical Pathology 2016 40 569–576.
 2858 (https://doi.org/10.1097/PAS.0000000000574)
- 2859 224 Morimoto R, Satoh F, Murakami O, Suzuki T, Abe T, Tanemoto M, Abe M, Uruno A,
 2860 Ishidoya S, Arai Y et al. Immunohistochemistry of a proliferation marker Ki67/MIB1 in
 2861 adrenocortical carcinomas: Ki67/ MIB1 labeling index is a predictor for recurrence of
 2862 adrenocortical carcinomas. Endocrine Journal 2008 55 49–55. (https://doi.
 2863 org/10.1507/endocrj.K07-079)
- 2864 225 Abiven G, Coste J, Groussin L, Anract P, Tissier F, Legmann P, Dousset B, Bertagna
 2865 X & Bertherat J. Clinical and biological features in the prognosis of adrenocortical cancer:
 2866 poor outcome of cortisol- secreting tumors in a series of 202 consecutive patients. Journal of
 2867 Clinical Endocrinology and Metabolism 2006 91 2650–2655.
- 2868 226 Burotto M, Tageja N, Rosenberg A, Mahalingam S, Quezado M, Velarde M, Edgerly
 2869 M & Fojo T. Brain metastasis in patients with adrenocortical carcinoma: a clinical series.
 2870 Journal of Clinical Endocrinology and Metabolism 2015 100 331–336. (https://doi.
 2871 org/10.1210/jc.2014-2650)
- 2872 Leboulleux S, Dromain C, Bonniaud G, Auperin A, Caillou B, Lumbroso J, Sigal R,
 2873 Baudin E & Schlumberger M. Diagnostic and prognostic value of 18-fluorodeoxyglucose
 2874 positron emission tomography in adrenocortical carcinoma: a prospective comparison with
 2875 computed tomography. Journal of Clinical Endocrinology and Metabolism 2006 91 920–925.
- 2876 228 Ardito A, Massaglia C, Pelosi E, Zaggia B, Basile V, Brambilla R, Vigna-Taglianti F,
 2877 Duregon E, Arena V, Perotti P et al. 18F-FDG PET/ CT in the post-operative monitoring of
 2878 patients with adrenocortical carcinoma. European Journal of Endocrinology 2015 173 749–
 2879 756. (https://doi.org/10.1530/EJE-15-0707)
- 2880 229 Berruti A, Fassnacht M, Baudin E, Hammer G, Haak H, Leboulleux S, Skogseid B,
 2881 Allolio B & Terzolo M. Adjuvant therapy in patients with adrenocortical carcinoma: a position
 2882 of an international panel. Journal of Clinical Oncology 2010 28 e401–e402; author reply
 2883 e403. (https://doi.org/10.1200/JCO.2009.27.5958)
- 2884 230 Megerle F, Herrmann W, Schloetelburg W, Ronchi CL, Pulzer A, Quinkler M,
 2885 Beuschlein F, Hahner S, Kroiss M & Fassnacht M. Mitotane monotherapy in patients with
 2886 advanced adrenocortical carcinoma. Journal of Clinical Endocrinology and Metabolism 2018
 2887 103 1686–1695. (https://doi.org/10.1210/jc.2017-02591)
- 2888 231 Volante M, Terzolo M, Fassnacht M, Rapa I, Germano A, Sbiera S, Daffara F,
 2889 Sperone P, Scagliotti G, Allolio B et al. Ribonucleotide reductase large subunit (RRM1) gene
 2890 expression may predict efficacy of adjuvant mitotane in adrenocortical cancer. Clinical
 2891 Cancer Research 2012 18 3452–3461. (https://doi.org/10.1158/1078-0432. CCR-11-2692)

- 2892 232 Ronchi CL, Sbiera S, Volante M, Steinhauer S, Scott-Wild V, Altieri B, Kroiss M, Bala
 2893 M, Papotti M, Deutschbein T et al. CYP2W1 is highly expressed in adrenal glands and is
 2894 positively associated with the response to mitotane in adrenocortical carcinoma. PLoS ONE
 2895 2014 9 e105855. (https://doi.org/10.1371/journal.pone.0105855)
- 2896 233 Terzolo M, Baudin AE, Ardito A, Kroiss M, Leboulleux S, Daffara F, Perotti P,
 2897 Feelders RA, deVries JH, Zaggia B et al. Mitotane levels predict the outcome of patients with
 2898 adrenocortical carcinoma treated adjuvantly following radical resection. European Journal of
 2899 Endocrinology 2013 169 263–270. (https://doi.org/10.1530/EJE-13-0242)
- 2900 234 Terzolo M, Daffara F, Ardito A, Zaggia B, Basile V, Ferrari L & Berruti A. Management
 2901 of adrenal cancer: a 2013 update. Journal of Endocrinological Investigation 2014 37 207–
 2902 217. (https://doi. org/10.1007/s40618-013-0049-2)
- 2903 235 Huang H & Fojo T. Adjuvant mitotane for adrenocortical cancer a recurring
 2904 controversy. Journal of Clinical Endocrinology and Metabolism 2008 93 3730–3732.
 2905 (https://doi.org/10.1210/jc.2008-0579)
- 2906 236 Terzolo M, Fassnacht M, Ciccone G, Allolio B & Berruti A. Adjuvant mitotane for
 2907 adrenocortical cancer working through uncertainty. Journal of Clinical Endocrinology and
 2908 Metabolism 2009 94 1879–1880. (https://doi.org/10.1210/jc.2009-0120)
- 2909 237 Polat B, Fassnacht M, Pfreundner L, Guckenberger M, Bratengeier K, Johanssen S,
 2910 Kenn W, Hahner S, Allolio B & Flentje M. Radiotherapy in adrenocortical carcinoma. Cancer
 2911 2009 115 2816–2823. (https:// doi.org/10.1002/cncr.24331)
- 238 Sabolch A, Else T, Jackson W, Williams A, Miller BS, Worden F, Hammer GD & Jolly
 2913 S. Improved local control with adjuvant radiation therapy in localized adrenocortical
 2914 carcinoma: a case- matched retrospective study. International Journal of Radiation Oncology
 2915 Biology Physics 2013 1 S84. (https://doi.org/10.1016/j. ijrobp.2013.06.219)
- 2916 239 Nelson DW, Chang SC, Bandera BC, Fischer TD, Wollman R & Goldfarb M. Adjuvant
 2917 radiation is associated with improved survival for select patients with non-metastatic
 2918 adrenocortical carcinoma. Annals of Surgical Oncology 2018 25 2060–2066. (https://doi.
 2919 org/10.1245/s10434-018-6510-x)
- 2920 240 Cerquetti L, Bucci B, Marchese R, Misiti S, De Paula U, Miceli R, Muleti A, Amendola
 2921 D, Piergrossi P, Brunetti E et al. Mitotane increases the radiotherapy inhibitory effect and
 2922 induces G2-arrest in combined treatment on both H295R and SW13 adrenocortical cell lines.
 2923 Endocrine-Related Cancer 2008 15 623–634. (https://doi. org/10.1677/erc.1.1315)
- 2924 241 Cerquetti L, Sampaoli C, Amendola D, Bucci B, Misiti S, Raza G, De Paula U,
 2925 Marchese R, Brunetti E, Toscano V et al. Mitotane sensitizes adrenocortical cancer cells to
 2926 ionizing radiations by involvement of the cyclin B1/CDK complex in G2 arrest and mismatch
 2927 repair enzymes modulation. International Journal of Oncology 2010 37 493–501.
- 242 Khan TS, Imam H, Juhlin C, Skogseid B, Grondal S, Tibblin S, Wilander E, Oberg K &
 2929 Eriksson B. Streptozocin and o,p'DDD in the treatment of adrenocortical cancer patients:
 2930 long-term survival in its adjuvant use. Annals of Oncology 2000 11 1281–1287. (https://doi.
 2931 org/10.1023/A:1008377915129)
- 243 Hermsen IGC, Gelderblom H, Kievit J, Romijn JA & Haak HR. Extremely long survival
 in six patients despite recurrent and metastatic adrenal carcinoma. European Journal of
 Endocrinology 2008 158 911–919. (https://doi.org/10.1530/EJE-07-0723)
- 2935 244 Bednarski BK, Habra MA, Phan A, Milton DR, Wood C, Vauthey N, Evans DB, Katz 2936 MH, Ng CS, Perrier ND et al. Borderline resectable adrenal cortical carcinoma: a potential

- role for preoperative chemotherapy. World Journal of Surgery 2014 38 1318–1327. (https://
 doi.org/10.1007/s00268-014-2484-4)
- 245 Wangberg B, Khorram-Manesh A, Jansson S, Nilsson B, Nilsson O, Jakobsson CE,
 Lindstedt S, Oden A & Ahlman H. The long-term survival in adrenocortical carcinoma with
 active surgical management and use of monitored mitotane. Endocrine-Related Cancer 2010
 17 265–272. (https://doi.org/10.1677/ERC-09-0190)
- 2943 246 Rangel C, Scattolin G, Pais-Costa SR, Vieira E & Gaio E. Neoadjuvant chemotherapy
 2944 and salvage surgery for an aldosterone-producing adrenal carcinoma with inferior vena cava
 2945 thrombus: case report and literature review. Asian Journal of Surgery 2013 36 134–136.
 2946 (https://doi.org/10.1016/j.asjsur.2012.08.008)
- 2947 Ronchi CL, Sbiera S, Kraus L, Wortmann S, Johanssen S, Adam P, Willenberg HS,
 2948 Hahner S, Allolio B & Fassnacht M. Expression of excision repair cross complementing
 2949 group 1 and prognosis in adrenocortical carcinoma patients treated with platinum-based
 2950 chemotherapy. Endocrine-Related Cancer 2009 16 907–918. (https:// doi.org/10.1677/ERC2951 08-0224)
- 248 Malandrino P, Al Ghuzlan A, Castaing M, Young J, Caillou B, Travagli JP, Elias D, de
 2953 Baere T, Dromain C, Paci A et al. Prognostic markers of survival after combined mitotane2954 and platinum-based chemotherapy in metastatic adrenocortical carcinoma. Endocrine2955 Related Cancer 2010 17 797–807. (https://doi.org/10.1677/ERC-09-0341)
- 2956 249 Roca E, Berruti A, Sbiera S, Rapa I, Oneda E, Sperone P, Ronchi CL, Ferrari L,
 2957 Grisanti S, Germano A et al. Topoisomerase2alpha and thymidylate synthase expression in
 2958 adrenocortical cancer. Endocrine- Related Cancer 2017 24 299–307.
 2959 (https://doi.org/10.1530/ERC-17-0095)
- 2960 250 Laufs V, Altieri B, Sbiera S, Kircher S, Steinhauer S, Beuschlein F, Quinkler M,
 2961 Willenberg HS, Rosenwald A, Fassnacht M et al. ERCC1 as predictive biomarker to
 2962 platinum-based chemotherapy in adrenocortical carcinomas. European Journal of
 2963 Endocrinology 2018 178 183–190. (https://doi.org/10.1530/EJE-17-0788)
- 2964 251 Bates SE, Shieh CY, Mickley LA, Dichek HL, Gazdar A, Loriaux DL & Fojo AT.
 2965 Mitotane enhances cytotoxicity of chemotherapy in cell lines expressing a multidrug
 2966 resistance gene (mdr-1/P-glycoprotein) which is also expressed by adrenocortical
 2967 carcinomas. Journal of Clinical Endocrinology and Metabolism 1991 73 18–29. (https://doi.
 2968 org/10.1210/jcem-73-1-18)
- 2969 252 Almeida MQ, Fragoso MC, Lotfi CF, Santos MG, Nishi MY, Costa MH, Lerario AM,
 2970 Maciel CC, Mattos GE, Jorge AA et al. Expression of IGF-II and its receptor in pediatric and
 2971 adult adrenocortical tumors. Journal of Clinical Endocrinology and Metabolism 2008 93
 2972 3524–3531. (https://doi.org/10.1210/jc.2008-0065)
- 2973 253 Boulle N, Logie A, Gicquel C, Perin L & Le Bouc Y. Increased levels of insulin-like
 2974 growth factor II (IGF-II) and IGF-binding protein-2 are associated with malignancy in sporadic
 2975 adrenocortical tumors. Journal of Clinical Endocrinology and Metabolism 1998 83 1713–
 2976 1720.
- 2977 254 Gicquel C, Bertagna X, Schneid H, Francillard-Leblond M, Luton JP, Girard F & Le
 2978 Bouc Y. Rearrangements at the 11p15 locus and overexpression of insulin-like growth factor2979 II gene in sporadic adrenocortical tumors. Journal of Clinical Endocrinology and Metabolism
 2980 1994 78 1444–1453.

- 2981 255 Giordano TJ, Thomas DG, Kuick R, Lizyness M, Misek DE, Smith AL, Sanders D,
 2982 Aljundi RT, Gauger PG, Thompson NW et al. Distinct transcriptional profiles of adrenocortical
 2983 tumors uncovered by DNA microarray analysis. American Journal of Pathology 2003 162
 2984 521–531. (https://doi.org/10.1016/S0002-9440(10)63846-1)
- 2985 256 Weber MM, Fottner C & Wolf E. The role of the insulin-like growth factor system in
 2986 adrenocortical tumourigenesis. European Journal of Clinical Investigation 2000 30
 2987 (Supplement 3) 69–75. (https://doi. org/10.1046/j.1365-2362.2000.0300s3069.x)
- 2988 257 Jones RL, Kim ES, Nava-Parada P, Alam S, Johnson FM, Stephens AW, Simantov 2989 R, Poondru S, Gedrich R, Lippman SM et al. Phase I study of intermittent oral dosing of the 2990 insulin-like growth factor-1 and insulin receptors inhibitor OSI-906 in patients with advanced 2991 solid tumors. Clinical Cancer Research 2015 21 693–700. (https://doi. org/10.1158/1078-2992 0432.CCR-14-0265)
- 2993 258 Naing A, Kurzrock R, Burger A, Gupta S, Lei X, Busaidy N, Hong D, Chen HX, Doyle
 2994 LA, Heilbrun LK et al. Phase I trial of cixutumumab combined with temsirolimus in patients
 2995 with advanced cancer. Clinical Cancer Research 2011 17 6052–6060. (https://doi.
 2996 org/10.1158/1078-0432.CCR-10-2979)
- 2997 259 Vezzosi D, Do Cao C, Hescot S, Bertherat J, Haissaguerre M, Bongard V, Drui D, De
 2998 La Fouchardiere C, Illouz F, Borson-Chazot F et al. Time until partial response in metastatic
 2999 adrenocortical carcinoma long-term survivors. Hormones and Cancer 2018 9 62–69.
 3000 (https://doi.org/10.1007/s12672-017-0313-6)
- Kroiss M, Quinkler M, Lutz WK, Allolio B & Fassnacht M. Drug interactions with
 mitotane by induction of CYP3A4 metabolism in the clinical management of adrenocortical
 carcinoma. Clinical Endocrinology 2011 75 585–591. (https://doi.org/10.1111/j.13652265.2011.04214.x)
- 3005 261 Faggiano A, Leboulleux S, Young J, Schlumberger M & Baudin E. Rapidly
 3006 progressing high o,p'DDD doses shorten the time required to reach the therapeutic threshold
 3007 with an acceptable tolerance: preliminary results. Clinical Endocrinology 2006 64 110–113.
 3008 (https:// doi.org/10.1111/j.1365-2265.2005.02403.x)
- Mauclere-Denost S, Leboulleux S, Borget I, Paci A, Young J, Al Ghuzlan A,
 Deandreis D, Drouard L, Tabarin A, Chanson P et al. High-dose mitotane strategy in
 adrenocortical carcinoma (ACC): prospective analysis of plasma mitotane measurement
 during the first three months of follow-up. European Journal of Endocrinology 2011 166 261–
 268. (https://doi.org/10.1530/EJE-11-0557)
- 3014 263 Terzolo M, Pia A, Berruti A, Osella G, Ali A, Carbone V, Testa E, Dogliotti L & Angeli
 3015 A. Low-dose monitored mitotane treatment achieves the therapeutic range with manageable
 3016 side effects in patients with adrenocortical cancer. Journal of Clinical Endocrinology and
 3017 Metabolism 2000 85 2234–2238.
- 3018 264 Terzolo M & Berruti A. Adjunctive treatment of adrenocortical carcinoma. Current
 3019 Opinion in Endocrinology, Diabetes and Obesity 2008 15 221–226.
 3020 (https://doi.org/10.1097/MED.0b013e3282fdf4c0)
- 3021 265 Kerkhofs TM, Baudin E, Terzolo M, Allolio B, Chadarevian R, Mueller HH, Skogseid
 3022 B, Leboulleux S, Mantero F, Haak HR et al. Comparison of two mitotane starting dose
 3023 regimens in patients with advanced adrenocortical carcinoma. Journal of Clinical
 3024 Endocrinology and Metabolism 2013 98 4759–4767. (https://doi.org/10.1210/ jc.2013-2281)

- 3025 266 Moolenaar AJ, van Slooten H, van Seters AP & Smeenk D. Blood levels of o,p'-DDD
 3026 following administration in various vehicles after a single dose and during long-term
 3027 treatment. Cancer Chemotherapy and Pharmacology 1981 7 51–54. (https://doi.org/10.1007/
 3028 BF00258213)
- 3029 267 Kerkhofs TM, Derijks LJ, Ettaieb MH, Eekhoff EM, Neef C, Gelderblom H, den Hartigh
 3030 J, Guchelaar HJ & Haak HR. Short-term variation in plasma mitotane levels confirms the
 3031 importance of trough level monitoring. European Journal of Endocrinology 2014 171 677–
 3032 683. (https://doi.org/10.1530/EJE-14-0388)
- 268 Daffara F, De Francia S, Reimondo G, Zaggia B, Aroasio E, Porpiglia F, Volante M,
 3034 Termine A, Di Carlo F, Dogliotti L et al. Prospective evaluation of mitotane toxicity in
 3035 adrenocortical cancer patients treated adjuvantly. Endocrine-Related Cancer 2008 15 1043–
 3036 1053. (https://doi.org/10.1677/ERC-08-0103)
- 3037 269 van Slooten H, Moolenaar AJ, van Seters AP & Smeenk D. The treatment of
 adrenocortical carcinoma with o,p'-DDD: prognostic implications of serum level monitoring.
 Buropean Journal of Cancer and Clinical Oncology 1984 20 47–53.
 (https://doi.org/10.1016/0277-5379(84)90033-6)
- Chortis V, Taylor AE, Schneider P, Tomlinson JW, Hughes BA, O'Neil DM, Libé R,
 Allolio B, Bertagna X, Bertherat J et al. Mitotane therapy in adrenocortical cancer induces
 CYP3A4 and inhibits 5α-reductase, explaining the need for personalized glucocorticoid and
 androgen replacement. Journal of Clinical Endocrinology and Metabolism 2013 98 161–171.
 (https://doi.org/10.1210/jc.2012-2851)
- Reimondo G, Puglisi S, Zaggia B, Basile V, Saba L, Perotti P, De Francia S, Volante
 M, Zatelli MC, Cannavo S et al. Effects of mitotane on the hypothalamic-pituitary-adrenal axis
 in patients with adrenocortical carcinoma. European Journal of Endocrinology 2017 177 361–
 367. (https://doi.org/10.1530/EJE-17-0452)
- 3050 272 Kerkhofs TM, Derijks LJ, Ettaieb H, den Hartigh J, Neef K, Gelderblom H, Guchelaar
 3051 HJ & Haak HR. Development of a pharmacokinetic model of mitotane: toward personalized
 3052 dosing in adrenocortical carcinoma. Therapeutic Drug Monitoring 2015 37 58–65.
 3053 (https://doi.org/10.1097/FTD.00000000000102)
- Alexandraki KI, Kaltsas GA, le Roux CW, Fassnacht M, Ajodha S, Christ-Crain M,
 Akker SA, Drake WM, WM, Edwards R, Allolio B et al. Assessment of serum-free cortisol
 levels in patients with adrenocortical carcinoma treated with mitotane: a pilot study. Clinical
 Endocrinology 2010 72 305–311. (https://doi.org/10.1111/ j.1365-2265.2009.03631.x)
- 3058 274 Russo M, Scollo C, Pellegriti G, Cotta OR, Squatrito S, Frasca F, Cannavo S & Gullo
 3059 D. Mitotane treatment in patients with adrenocortical cancer causes central hypothyroidism.
 3060 Clinical Endocrinology 2016 84 614–619. (https://doi.org/10.1111/cen.12868)
- Tada H, Nohara A, Kawashiri MA, Inazu A, Mabuchi H & Yamagishi M. Marked
 transient hypercholesterolemia caused by low-dose mitotane as adjuvant chemotherapy for
 adrenocortical carcinoma. Journal of Atherosclerosis and Thrombosis 2014 21 1326–1329.
 (https://doi.org/10.5551/jat.27557)
- 3065 276 Nieman LK, Biller BM, Findling JW, Murad MH, Newell-Price J, Savage MO & Tabarin
 3066 A. Treatment of Cushing's syndrome: an Endocrine Society Clinical Practice Guideline.
 3067 Journal of Clinical Endocrinology and Metabolism 2015 100 2807–2831. (https://doi.
- 3068 org/10.1210/jc.2015-1818)

- Claps M, Cerri S, Grisanti S, Lazzari B, Ferrari V, Roca E, Perotti P, Terzolo M,
 Sigala S & Berruti A. Adding metyrapone to chemotherapy plus mitotane for Cushing's
 syndrome due to advanced adrenocortical carcinoma. Endocrine 2017 61 169–172.
 (https://doi. org/10.1007/s12020-017-1428-9)
- 278 Castinetti F, Fassnacht M, Johanssen S, Terzolo M, Bouchard P, Chanson P, Do Cao
 3074 C, Morange I, Pico A, Ouzounian S et al. Merits and pitfalls of mifepristone in Cushing's
 3075 syndrome. European Journal of Endocrinology 2009 160 1003–1010.
- 3076 (https://doi.org/10.1530/EJE-09-0098)
- Chow E, Hoskin P, Mitera G, Zeng L, Lutz S, Roos D, Hahn C, van der Linden Y,
 Hartsell W & Kumar E. Update of the international consensus on palliative radiotherapy
 endpoints for future clinical trials in bone metastases. International Journal of Radiation
 Oncology, Biology, Physics 2012 82 1730–1737. (https://doi.org/10.1016/j.
 ijrobp.2011.02.008)
- 280 Pin Y, Paix A, Le Fevre C, Antoni D, Blondet C & Noel G. A systematic review of
 palliative bone radiotherapy based on pain relief and retreatment rates. Critical Reviews in
 Oncology/ Hematology 2018 123 132–137. (https://doi.org/10.1016/j. critrevonc.2018.01.006)
- 3085 281 Ferrell BR, Temel JS, Temin S, Alesi ER, Balboni TA, Basch EM, Firn JI, Paice JA,
 3086 Peppercorn JM, Phillips T et al. Integration of palliative care into standard oncology care:
 3087 American Society of Clinical Oncology Clinical Practice Guideline update. Journal of Clinical
 3088 Oncology 2017 35 96–112. (https://doi.org/10.1200/JCO.2016.70.1474)
- 3089 282 Herrmann LJ, Heinze B, Fassnacht M, Willenberg HS, Quinkler M, Reisch N, Zink M,
 3090 Allolio B & Hahner S. TP53 germline mutations in adult patients with adrenocortical
 3091 carcinoma. Journal of Clinical Endocrinology and Metabolism 2012 97 E476–E485.
 3092 (https://doi. org/10.1210/jc.2011-1982)
- Raymond VM, Else T, Everett JN, Long JM, Gruber SB & Hammer GD. Prevalence of
 germline TP53 mutations in a prospective series of unselected patients with adrenocortical
 carcinoma. Journal of Clinical Endocrinology and Metabolism 2013 98 E119–E125.
 (https://doi. org/10.1210/jc.2012-2198)
- 3097 284 Waldmann J, Patsalis N, Fendrich V, Langer P, Saeger W, Chaloupka B,
 3098 Ramaswamy A, Fassnacht M, Bartsch DK & Slater EP. Clinical impact of TP53 alterations in
 3099 adrenocortical carcinomas. Langenbecks Archives of Surgery 2012 397 209–216.
 3100 (https://doi. org/10.1007/s00423-011-0868-6)
- Raymond VM, Everett JN, Furtado LV, Gustafson SL, Jungbluth CR, Gruber SB,
 Hammer GD, Stoffel EM, Greenson JK, Giordano TJ et al. Adrenocortical carcinoma is a
 lynch syndrome-associated cancer. Journal of Clinical Oncology 2013 31 3012–3018.
 (https://doi. org/10.1200/JCO.2012.48.0988)
- 3105 286 Stoffel EM, Mangu PB, Gruber SB, Hamilton SR, Kalady MF, Lau MW, Lu KH, Roach
 3106 N & Limburg PJ. Hereditary colorectal cancer syndromes: American Society of Clinical
 3107 Oncology Clinical Practice Guideline endorsement of the familial risk-colorectal cancer:
 3108 European Society for Medical Oncology Clinical Practice Guidelines. Journal of Clinical
 3109 Oncology 2015 33 209–217. (https://doi. org/10.1200/JCO.2014.58.1322)
- 287 Daly MB, Pilarski R, Berry M, Buys SS, Farmer M, Friedman S, Garber JE, Kauff ND,
 3111 Khan S, Klein C et al. NCCN Guidelines Insights: genetic/familial high-risk assessment:
 3112 breast and ovarian, version 2.2017. Journal of the National Comprehensive Cancer Network
 3113 2017 15 9–20. (https://doi.org/10.6004/jnccn.2017.0003)

- 3114 288 Kratz CP, Achatz MI, Brugieres L, Frebourg T, Garber JE, Greer MC, Hansford JR,
 3115 Janeway KA, Kohlmann WK, McGee R et al. Cancer screening recommendations for
 3116 individuals with Li-Fraumeni syndrome. Clinical Cancer Research 2017 23 e38–e45.
 3117 (https://doi. org/10.1158/1078-0432.CCR-17-0408)
- Ballinger ML, Best A, Mai PL, Khincha PP, Loud JT, Peters JA, Achatz MI, Chojniak
 R, Balieiro da Costa A, Santiago KM et al. Baseline surveillance in Li-Fraumeni syndrome
 using whole-body magnetic resonance imaging: a meta-analysis. JAMA Oncology 2017 3
 1634–1639. (https://doi.org/10.1001/jamaoncol.2017.1968)
- 3122 290 Gupta S, Provenzale D, Regenbogen SE, Hampel H, Slavin TP Jr, Hall MJ, Llor X,
 3123 Chung DC, Ahnen DJ, Bray T et al. NCCN guidelines insights: genetic/familial high-risk
 3124 assessment: colorectal, version 3.2017. Journal of the National Comprehensive Cancer
 3125 Network 2017 15 1465–1475. (https://doi.org/10.6004/jnccn.2017.0176)
- 3126 291 Stoffel EM, Mangu PB & Limburg PJ. Hereditary colorectal cancer syndromes:
 3127 American Society of Clinical Oncology clinical practice guideline endorsement of the familial
 3128 risk-colorectal cancer: European Society for Medical Oncology clinical practice guidelines.
- 3129 Journal of Oncology Practice 2015 11 e437–e441. (https://doi.
- 3130 org/10.1200/JOP.2015.003665)
- 3131 292 Else T. Association of adrenocortical carcinoma with familial cancer susceptibility
- 3132 syndromes. Molecular and Cellular Endocrinology 2012 351 66–70.
- 3133 (https://doi.org/10.1016/j.mce.2011.12.008)
- McDonnell CM & Zacharin MR. Adrenal cortical tumours: 25 years' experience at the
 Royal Children's Hospital, Melbourne. Journal of Paediatrics and Child Health 2003 39 682–
 685. (https://doi. org/10.1046/j.1440-1754.2003.00268.x)
- 294 Custodio G, Parise GA, Kiesel Filho N, Komechen H, Sabbaga CC, Rosati R, Grisa L,
 Parise IZ, Pianovski MA, Fiori CM et al. Impact of neonatal screening and surveillance for the
 TP53 R337H mutation on early detection of childhood adrenocortical tumors. Journal of
 Clinical Oncology 2013 31 2619–2626. (https://doi.org/10.1200/ JCO.2012.46.3711)
- Wasserman JD, Novokmet A, Eichler-Jonsson C, Ribeiro RC, Rodriguez-Galindo C,
 Zambetti GP & Malkin D. Prevalence and functional consequence of TP53 mutations in
 pediatric adrenocortical carcinoma: a children's oncology group study. Journal of Clinical
 Oncology 2015 33 602–609. (https://doi.org/10.1200/ JCO.2013.52.6863)
- 3145 296 Eschler DC, Kogekar N & Pessah-Pollack R. Management of adrenal tumors in
 3146 pregnancy. Endocrinology Metabolism Clinics of North America 2015 44 381–397.
 3147 (https://doi.org/10.1016/j. ecl.2015.02.006)
- Abiven-Lepage G, Coste J, Tissier F, Groussin L, Billaud L, Dousset B, Goffinet F,
 Bertagna X, Bertherat J & Raffin-Sanson ML. Adrenocortical carcinoma and pregnancy:
 clinical and biologicalfeatures and prognosis. European Journal of Endocrinology 2010 163
 793–800. (https://doi.org/10.1530/EJE-10-0412)
- Sirianni R, Zolea F, Chimento A, Ruggiero C, Cerquetti L, Fallo F, Pilon C, Arnaldi G,
 Carpinelli G, Stigliano A et al. Targeting estrogen receptor-alpha reduces adrenocortical
 cancer (ACC) cell growth in vitro and in vivo: potential therapeutic role of selective estrogen
 receptor modulators (SERMs) for ACC treatment. Journal of Clinical Endocrinology and
 Metabolism 2012 97 E2238–E2250. (https://doi. org/10.1210/jc.2012-2374)

Tripto-Shkolnik L, Blumenfeld Z, Bronshtein M, Salmon A & Jaffe A. Pregnancy in a
patient with adrenal carcinoma treated with mitotane: a case report and review of literature.
Journal of Clinical Endocrinology and Metabolism 2013 98 443–447.

300 de Corbiere P, Ritzel K, Cazabat L, Ropers J, Schott M, Libe R, Koschker AC,
Leboulleux S, Deutschbein T, Do Cao C et al. Pregnancy in women previously treated for an
adrenocortical carcinoma. Journal of Clinical Endocrinology and Metabolism 2015 100 4604–
4611. (https://doi.org/10.1210/jc.2015-2341)

- 301 Hescot S, Seck A, Guerin M, Cockenpot F, Huby T, Broutin S, Young J, Paci A,
 Baudin E & Lombes M. Lipoprotein-free mitotane exerts high cytotoxic activity in
 adrenocortical carcinoma. Journal of Clinical Endocrinology and Metabolism 2015 100 2890–
 2898. (https://doi. org/10.1210/JC.2015-2080)
- 302 Hescot S, Slama A, Lombes A, Paci A, Remy H, Leboulleux S, Chadarevian R,
 Trabado S, Amazit L, Young J et al. Mitotane alters mitochondrial respiratory chain activity by
 inducing cytochrome c oxidase defect in human adrenocortical cells. Endocrine-Related
 Cancer 2013 20 371–381. (https://doi.org/10.1530/ERC-12-0368)
- 303 Sbiera S, Leich E, Liebisch G, Sbiera I, Schirbel A, Wiemer L, Matysik S, Eckhardt C,
 3173 Gardill F, Gehl A et al. Mitotane inhibits sterol-O-acyl transferase 1 triggering lipid-mediated
 and apoptosis in adrenocortical carcinoma cells. Endocrinology
 3175 2015 156 3895–3908. (https://doi.org/10.1210/en.2015-1367)
- 304 Hescot S, Amazit L, Lhomme M, Travers S, DuBow A, Battini S, Boulate G, Namer IJ,
 Lombes A, Kontush A et al. Identifying mitotane-induced mitochondria-associated
 membranes dysfunctions: metabolomic and lipidomic approaches. Oncotarget 2017 8
 109924–109940.
- 305 Arlt W, Biehl M, Taylor AE, Hahner S, Libe R, Hughes BA, Schneider P, Smith DJ,
 3181 Stiekema H, Krone N et al. Urine steroid metabolomics as a biomarker tool for detecting
 3182 malignancy in adrenal tumors. Journal of Clinical Endocrinology and Metabolism 2011 96
 3183 3775–3784. (https:// doi.org/10.1210/jc.2011-1565)
- 306 Kerkhofs TM, Kerstens MN, Kema IP, Willems TP & Haak HR. Diagnostic value of
 urinary steroid profiling in the evaluation of adrenal tumors. Hormones and Cancer 2015 6
 168–175. (https://doi. org/10.1007/s12672-015-0224-3)
- 307 Taylor DR, Ghataore L, Couchman L, Vincent RP, Whitelaw B, Lewis D, Diaz-Cano
 S, Galata G, Schulte KM, Aylwin S et al. A 13-steroid serum panel based on LC-MS/MS: use
 in detection of adrenocortical carcinoma. Cliniccal Chemistry 2017 63 1836–1846.
 (https://doi.org/10.1373/clinchem.2017.277624)
- 3191 308 Hines JM, Bancos I, Bancos C, Singh RD, Avula AV, Young WF, Grebe SK & Singh
 3192 RJ. High-resolution, accurate-mass (HRAM) mass spectrometry urine steroid profiling in the
 3193 diagnosis of adrenal disorders. Clinical Chemistry 2017 63 1824–1835. (https://doi.
 3194 org/10.1373/clinchem.2017.271106)
- 309 Pinzani P, Scatena C, Salvianti F, Corsini E, Canu L, Poli G, Paglierani M, Piccini V,
 Pazzagli M, Nesi G et al. Detection of circulating tumor cells in patients with adrenocortical
 carcinoma: a monocentric preliminary study. Journal of Clinical Endocrinology and
 Metabolism 2013 98 3731–3738. (https://doi.org/10.1210/jc.2013-1396)
- 3199 310 Chabre O, Libe R, Assie G, Barreau O, Bertherat J, Bertagna X, Feige JJ & Cherradi
 3200 N. Serum miR-483-5p and miR-195 are predictive of recurrence risk in adrenocortical cancer
 3201 patients. Endocrine-Related Cancer 2013 20 579–594.

3202 311 Szabo DR, Luconi M, Szabo PM, Toth M, Szucs N, Horanyi J, Nagy Z, Mannelli M,
3203 Patocs A, Racz K et al. Analysis of circulating microRNAs in adrenocortical tumors.
3204 Laboratory Investigation 2014 94 331–339. (https://doi.org/10.1038/labinvest.2013.148)

3205 312 Perge P, Butz H, Pezzani R, Bancos I, Nagy Z, Paloczi K, Nyiro G, Decmann A, Pap
3206 E, Luconi M et al. Evaluation and diagnostic potential of circulating extracellular vesicle3207 associated microRNAs in adrenocortical tumors. Scientific Reports 2017 7 5474. (https://doi.
3208 org/10.1038/s41598-017-05777-0)

3209 313 Creemers SG, Korpershoek E, Atmodimedjo PN, Dinjens WNM, van Koetsveld PM, 3210 Feelders RA & Hofland LJ. Identification of mutations in cell-free circulating tumor DNA in 3211 adrenocortical carcinoma: a case series. Journal of Clinical Endocrinology and Metabolism 3212 2017 102 3611–3615. (https://doi.org/10.1210/jc.2017-00174)

- 3213 314 Garinet S, Nectoux J, Neou M, Pasmant E, Jouinot A, Sibony M, Orhant L, Pipoli da
 3214 Fonseca J, Perlemoine K, Bricaire L et al. Detection and monitoring of circulating tumor DNA
- in adrenocortical carcinoma. Endocrine-Related Cancer 2018 25 L13–L17.
- 3216 (https://doi.org/10.1530/ERC-17-0467)

3217