Transcriptome in paraffin samples for the diagnosis and prognosis of adrenocortical carcinoma

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Abstract

Design: Molecular classification is important for the diagnosis and prognosis of adrenocortical tumors (ACT). Transcriptome profiles separate adrenocortical adenomas ‘C2’ from carcinomas, and identify two groups of carcinomas ‘C1A’ and ‘C1B’, of poor and better prognosis respectively. However, many ACT cannot be profiled because of improper or absent freezing procedures, a mandatory requirement so far. The main aim was to determine transcriptome profiles on formalin-fixed paraffin-embedded (FFPE) samples, using the new 3’-end RNA-sequencing technology. A secondary aim was to demonstrate the ability of this technique to explore large FFPE archives, by focusing on the rare oncocytic ACT variants.

Methods: We included 131 ACT: a training cohort from Cochin hospital and an independent validation cohort from Wuerzburg hospital. The 3’ transcriptome was generated from FFPE samples using QuantSeq (Lexogen, Vienna, Austria) and NextSeq500 (Illumina, San Diego, CA, USA).

Results: In the training cohort, unsupervised clustering identified three groups: ‘C1A’ aggressive carcinomas (n = 28, 29%), ‘C1B’ more indolent carcinomas (n = 28, 29%), and ‘C2’ adenomas (n = 39, 41%). The prognostic value of FFPE transcriptome was confirmed in the validation cohort (5-year OS: 26% in ‘C1A’ (n = 26) and 100% in ‘C1B’ (n = 10), P = 0.003). FFPE transcriptome was an independent prognostic factor in a multivariable model including tumor stage and Ki-67 (OS HR: 7.5, P = 0.01). Oncocytic ACT (n = 19) did not form any specific cluster. Oncocytic carcinomas (n = 6) and oncocytic ACT of uncertain malignant potential (n = 4) were all in ‘C1B’.

Conclusions: The 3’ RNA-sequencing represents a convenient solution for determining ACT molecular class from FFPE samples. This technique should facilitate routine use and large retrospective studies.
Introduction

Adrenocortical carcinoma (ACC) is a rare endocrine cancer (1–2 per million per year) with an overall poor prognosis (1). Diagnosis is critical for patient management, but difficult owing to the rarity of the disease. Indeed, ACC is rare among adrenocortical tumors (ACT), the majority being adrenocortical adenomas (ACA). A pathology malignancy score called the Weiss score is most commonly used for diagnosis (2, 3). However, tumors with intermediate scores and rare histologic variants cannot be easily classified. In particular, determining the malignancy of oncocytic variants, composed of large cells with eosinophilic granulations, is challenging (4, 5).

The 5-year overall survival of ACC is generally below 40%, but heterogeneous (6). Prognostic stratification is important to guide treatment decision, especially in the adjuvant setting (7). The main prognostic factors used in clinical practice are tumor stage (8) and Ki-67 proliferation index (9). Other prognostic factors commonly include cortisol secretion (10) and age (11).

Previous genomic studies have demonstrated a convergence of genomic alterations into distinct molecular subtypes (12, 13). Transcriptome profiles represent the cornerstone of molecular classification. Transcriptome profiles separate adrenocortical adenomas (C2) from carcinomas and identify two groups of ACC strongly associated with outcome: the ‘C1A’ subgroup, with an overexpression of proliferative genes and a poor prognosis, and the ‘C1B’ subgroup, with an immune signature and a better prognosis (12, 13, 14, 15). Specific mRNA markers, such as the differential gene expression of BUB1B-PINK1, have then been deduced from transcriptome signatures. These targeted markers improve diagnosis and prognostic assessment of ACC (16, 17, 18). However, transcriptome profiles and their targeted surrogate markers are up to now limited to high-quality tumor RNA, requiring high-quality frozen tissue samples. Obtaining this type of material is demanding in routine practice, and often not achieved for ACC patients.

The recent 3’ RNA-sequencing technology captures and sequences short fragments at the 3’ end of polyadenylated RNA, which are less prone to RNA degradation. Unlike gene expression microarrays or standard full-length RNA-sequencing, this technique requires only a low amount and low-quality RNA, compatible with formalin-fixed paraffin-embedded (FFPE) samples (19).

In this study, we performed 3’ RNA-sequencing on FFPE samples of adrenocortical tumors. We evaluated the diagnostic and prognostic value of FFPE transcriptome classification, investigated the transcriptome classification of oncocytic variants, and assessed intra-tumor heterogeneity.

Subjects and methods

Patients

A training cohort of 95 ACT patients from Cochin Hospital (Paris, France) was used to explore FFPE transcriptome profiles of classical ACC, ACA and oncocytic ACT. All consecutive primary tumor specimens of classical ACC and oncocytic ACT collected between 2008 and 2018 were included. A control group of classical ACA was randomly selected over the same period. A fragment of tumor tissue was frozen in liquid nitrogen immediately after surgery and the remaining tumor was fixed in formalin and embedded in paraffin. The diagnosis of malignancy was established by an experienced endocrine pathologist (MS and FV) using the Weiss score (2, 3) (ACC if ≥3 malignancy criteria) or the Lin–Weiss–Bisceglia score (4) for oncocytic tumors (ACC if ≥1 major criteria, ACT of undetermined malignant potential if ≥1 minor criteria). For eight patients, two different regions of the primary tumor were sampled to assess the intratumor heterogeneity of ACT transcriptome. For 5/8 tumors, distinct nuclear and cytologic features were observed between the two regions.

A validation cohort of 36 ACC patients from Wuerzburg University Hospital (Wuerzburg, Germany) was then used to test the prognostic value of FFPE transcriptome profiles.

Clinical information was obtained from the European Network for the Study of Adrenal Tumors (ENSA T) registry (20) (Supplementary Table 1, see section on supplementary materials given at the end of this article).

Written consent has been obtained from each patient or subject after a full explanation of the purpose and nature of all procedures used. The study was conducted in accordance with the Declaration of Helsinki and approved by the Comité de protection des personnes Ile de France 1 (application no. 13311) and the Ethics Committee of the University of Wuerzburg (registration no. 88/11).

Targeted PCR marker on frozen samples

The differential gene expression of BUB1B-PINK1, recapitulating the ‘C1A’ and ‘C1B’ transcriptome difference, was used to predict ‘poor-outcome’ and ‘better-outcome’ ACC, as previously published (15, 16).
3’ RNA-seqencing on FFPE samples

FFPE material was selected by an experienced endocrine pathologist. RNA extraction was performed from 10 × 6 µm FFPE tissue sections using the Maxwell 16 LEV RNA FFPE kit and a Maxwell 16 Instrument (Promega). RNA was quantified using a Qubit fluorometer (Thermo Fisher Scientific) and analyzed by electrophoresis on a Bioanalyzer 2100 (Agilent Technologies). As expected, FFPE-extracted RNA showed degraded profiles, with RNA Integrity Number <3. No selection based on RNA quality was carried out.

Sequencing libraries were prepared at the Genomics Platform of the Cochin Institute, following the QuantSeq 3’ mRNA-Seq protocol (Lexogen, Vienna, Austria), starting from 100 to 500 ng RNA. Single read sequencing (1 ×75bp) was performed on a NextSeq 550 platform (Illumina, San Diego, CA, USA).

Transcriptome analyses

FASTQ sequences were aligned on GRCh38 human reference genome with STAR(v2.7.3a) (21). Read counts (Supplementary Table 2) were normalized with DESeq2 (v1.28.1) (22). Genes were filtered by s.d. to select the 2000 most variable genes for subsequent analyses (classification, principal component analysis, and transcriptome prediction).

Unsupervised classification of ACT transcriptome profiles was performed in the 95 patients of the Cochin cohort. Among tumors with two distinct regions, only the most histologically aggressive region was included in unsupervised classification. Groups with similar gene expression profiles were identified using non-negative matrix factorization (NMF) consensus clustering with 200 iterations, implemented in the NMF R package (v0.22.0) (23). The best rank of factorization (k = 3) was chosen from both cophenetic and average silhouette width score profiles. Adrenal differentiation, proliferation and immune signatures (Supplementary Table 3) were previously proposed to characterize transcriptome subgroups in the ACC TCGA study (13). The overall level of activation of these signatures was calculated in each sample using a single sample GSEA implemented in GSVA R package (v1.36.2) (24). Differentially expressed genes between NMF groups were identified using the Kruskal–Wallis test, with Benjamini-Hochberg correction for multiple testing. Gene-set enrichment analyses were performed with the GSEA pre-ranked method (v4.0.3) (25), using the ‘Hallmarks’ annotation v7.1 (MSigDB).

Principal component analysis (PCA) was performed to visualize individuals in the two-dimension views that capture the greatest amount of variability, using factoextra (v1.0.6) and FactoMineR (v2.3) R packages (26). The 95 unique ACT from Cochin were used to compute PCA. ACT with two distinct tumor regions and ACC from the validation cohort was then projected as supplementary individuals on the same representation.

Finally, a transcriptome-based model was developed to classify any new ACC sample into C1A/C1B prognostic groups. Read counts were normalized on the sequencing depth only using the formula:

\[
\log_2(1 + \text{CPM}), \text{ with CPM : counts per million}
\]

With this normalization, additional samples can be processed independently from the original cohort. A ridge penalized regression with ten-fold cross-validation was applied to predict the C1A/C1B status on the training (Cochin) cohort, using the glmnet R package (v4.0-2) (27). This model was then applied to the validation (Wuerzburg) cohort.

Statistical analysis

Comparisons between groups were performed with the Wilcoxon and Kruskal–Wallis tests for quantitative variables, and with Fisher’s test for qualitative variables.

Disease-free survival (DFS) was analyzed in patients with ENSAT stage I–III ACC and was defined as the time from primary tumor resection until recurrence or last follow-up. Overall survival (OS) was analyzed in stage I–IV ACC, and defined as the time from pathological diagnosis until death or last follow-up. Survival curves were obtained with Kaplan–Meier estimates and compared with a log-rank test. Cox proportional hazards regression was used to identify variables associated with DFS and OS. The proportional hazards assumption was checked for each model using graphical methods based on Kaplan–Meier curves and the scaled Schoenfield residuals. Significant variables were combined into multivariable models.

All P-values were two-sided, and the level of significance was set at P < 0.05. Calculations were performed using R statistical software (v4.1.1, R Stats, survival and survcomp packages).

Results

Patient characteristics

A training cohort of 95 ACT patients from Cochin hospital (43 classical ACC, 33 classical ACA and 19 oncocytic ACT), and an independent validation cohort of 36 ACC
patients from Wuerzburg hospital were included (Table 1 and Supplementary Table 1). Median follow-up were 40.3 months (95% CI: 2.1–136.1) in the training cohort and 30.5 months (95 CI: 4.0–225.9) in the validation cohort.

**Impact of sample age on sequencing data**

The median Q30 per sample was 88%, ranging from 84–91%. A median of 11 665 648 aligned reads per sample (range 3 255 802–22 861 798) were obtained. Sample age may impact both the amount (median number of aligned reads 12 780 317 in the 10 oldest samples vs 16 853 879 in the 10 most recent samples, \( P = 0.001 \)) and quality of reads (median Q30 87% in the 10 oldest samples vs 90% in the 10 most recent samples, \( P < 10^{-4} \)). In spite of slightly lower quality, transcriptome profiles were not impacted, since no association was observed between transcriptome classification and sample age (data not shown). Moreover, no technical failures were observed, even for samples collected 10 years ago.

**FFPE transcriptome classification of ACT**

In the training cohort, unsupervised consensus clustering of ACT transcriptome identified three distinct groups, closely associated with clinical, pathological and molecular characteristics (Fig. 1): (i) a group of aggressive ACC, enriched in advanced, high-grade tumors, showing high rates of recurrence (16/28, 57%) and death (14/28, 50%). This group was classified as poor-outcome with the targeted PCR marker BUB1B-PINK1; (ii) a group of more indolent ACC, enriched in localized, low-grade tumors, showing low rates of recurrence (4/28, 14%) and death (3/28, 11%). This group was classified as better-outcome with the targeted PCR marker BUB1B-PINK1; (iii) a group of ACA.

In terms of gene expression profiles, these FFPE transcriptome groups correspond to the known ACT classes (12, 13, 15) (Fig. 1A, B, C, D, E and Supplementary Tables 4, 5), with (i) high expression of cell-cycle genes such as TOP2A, MKI67 or BUB1 and enrichment in G2M_CHECKPOINT and E2F_TARGETS gene sets in the aggressive ACC group, previously referred to as the ‘CIA’ transcriptome class; (ii) an immune signature, with enrichment in ALLOGRAFT_REJECTION, INTERFERON_GAMMA_RESPONSE and IL6_JAK_STAT3_SIGNALING gene sets in the more indolent ACC group, previously referred to as the ‘C1B’ transcriptome class; (iii) high adrenal differentiation scores, and high expression of steroid enzymes in the ACA group, previously referred to as the ‘C2’ transcriptome class.

**Table 1** Patients’ characteristics. Values are expressed in \( n \) (%) for qualitative variables and in median (range) for quantitative variables. \( P \)-values < 0.05 are indicated in bold.

|                      | Training cohort | Validation cohort | \( P \)-value * |
|----------------------|-----------------|-------------------|----------------|
| \( n \)               | 42              | 49 + 4*           | 0.61           |
| Age (years)          | 51 (25–85)      | 47 (21–73)        | 0.34           |
| Sex                  | Male            | Female            | 0.33           |
| Female               | 33 (79%)        | 41 (77%)          | 0.46           |
| Male                 | 21 (9%)         | 12 (23%)          |                |
| Cortisol secretion   | Yes             | No                | 0.82           |
| Yes                  | 32 (78%)        | 33 (62%)          | 0.005          |
| No                   | 9 (22%)         | 20 (38%)          |                |
| Not available        | 1               |                   |                |
| Subtype              | Classical       | Oncytic           |                |
| Classical            | 33 (79%)        | 43 (81%)          | 0.0001         |
| Oncytic              | 9 (21%)         | 10 (19%)          |                |
| Not available        | 2               |                   |                |
| Weiss score          |                 |                   |                |
| 0–1                  | 26 (62%)        |                   | 0.17           |
| 2                    | 13 (31%)        |                   |                |
| 3                    | 3 (7%)          | 8 (15%)           |                |
| 4–6                  | 17 (32%)        | 14 (47%)          |                |
| 7–9                  | 28 (53%)        | 15 (50%)          |                |
| Ki-67                |                 |                   | 0.008          |
| <10%                 | 29 (100%)       | 23 (49%)          | 0.17           |
| 10–19%               | 0 (0%)          | 7 (15%)           |                |
| ≥20%                 | 0 (0%)          | 17 (36%)          |                |
| Not available        | 13              | 6                  |                |
| Relapse or metastases|                 |                   | <10^{-4}       |
| No                   | 34 (100%)       | 33 (62%)          |                |
| Yes                  | 0 (0%)          | 20 (38%)          |                |
| Not available        | 8               | 30 (83%)          |                |
| Death                |                 |                   | 0.27           |
| No                   | 34 (100%)       | 36 (68%)          |                |
| Yes                  | 0 (0%)          | 17 (32%)          |                |
| Not available        | 8               | 16 (44%)          |                |

*ACC \( n = 49 \) and ACT of uncertain malignant potential \( n = 4 \); \( \dagger \) \( P \)-values are provided for the comparison between ACC and ACT of uncertain malignant potential from the training cohort and ACC from the validation cohort.

**FFPE transcriptome profiles of oncocytic ACT**

In this cohort, 19/95 ACT were oncocytic tumors, including 6 ACC, 4 ACT of uncertain malignant potential and 9 ACA according to the Lin–Weiss–Bisceglia classification (4). In unsupervised transcriptome classification, these ACT did
Figure 1
FFPE transcriptome classification of ACT in the training cohort. (A) Unsupervised classification of ACT based on the 2000 most variable genes identifies three main groups, corresponding to 'C1A' aggressive carcinomas, 'C1B' indolent carcinomas and 'C2' adenomas. (B) Heatmap of the three non-negative matrix factorization (NMF) ranks used for generating the unsupervised classification. (C) Expression of steroid (adrenal differentiation), proliferation and immune signatures in each ACT. (D) Expression profiles of the top 50 most significantly different genes among the three transcriptome groups. (E) Consensus matrix representing the similarity between samples over iterative clustering algorithms. Rows and columns are patient samples. Consensus matrix values range from 0 (never clustered together) to 1 (always clustered together).
not segregate into any specific cluster of oncocytic tumors. Instead, oncocytic ACT scattered, evenly mixed with other ACT within C2 adenomas and C1B carcinomas (Fig. 1A), with no specific transcriptome signature (Supplementary Table 6). In particular, all oncocytic ACC and oncocytic ACT of uncertain malignant potential clustered with C1B carcinomas.

**FFPE transcriptome profiles in different tumor regions**

Transcriptome heterogeneity was explored in two different regions of the primary tumor for eight patients (Fig. 2A and B). For all patients, the two samples were concordant in terms of transcriptome classification, in spite of distinct nuclear and cytological features for 5/8 tumors (Fig. 2C). For these patients, tumor regions with mainly ‘benign’ features on pathology were properly classified as carcinomas (C1A or C1B), following the transcriptome class of their most aggressive counterpart (Fig. 2C).

**C1A and C1B carcinomas showed a distinct outcome (Supplementary Fig. 1), as previously established (12, 13, 15, 16). FFPE transcriptome prognostic value through proper C1A/C1B classification was confirmed in an independent validation cohort. Transcriptome profiles of these independent samples were close to either C1A (n=26) or C1B (n=10) carcinomas from the training cohort (Fig. 3A). We next developed a predictive model of transcriptome class, using cross-validated ridge regression, for determining the prognosis of individual patients. This model was optimized to discriminate C1A from C1B carcinomas on the training cohort. On the validation cohort, predicted C1A carcinomas were associated with shorter DFS (median 7.5 vs 240 months, P = 0.0014) and OS (median 30 months vs not reached, P = 0.0027) compared to predicted C1B carcinomas (Fig. 3B and C). Of note, 5-year OS was 26% (95 CI 12 to 56%) in predicted C1A and 100% in predicted C1B carcinomas.

**Multivariate prognostic analyses**

The prognostic value of FFPE transcriptome classification was then tested against known ACC prognostic factors in the whole ACC cohort. C1A cluster was associated with cortisol-secreting tumors (P = 0.001), high Ki-67 (P < 10^{-2}) and incomplete surgical resection (P = 0.0002) (Supplementary Table 7).

In stage I-III ACC, ENSAT stage, Ki-67 index and molecular class were identified as independent prognostic factors of DFS (Table 2). In stage I-IV ACC, ENSAT stage and molecular class were the only independent prognostic factors of OS (Table 3).

**Discussion**

This study shows the feasibility of ACT transcriptome from FFPE samples. All samples were suitable for the technique, even those collected more than 10 years ago. We showed that diagnostic and prognostic transcriptome classification of ACT can be obtained from FFPE samples using 3' RNA-sequencing. We were able to assess the transcriptome classification of oncocytic ACT and the intratumor heterogeneity of ACT transcriptome and to develop a new transcriptome-based model for ACC prognostication in clinical routine.

Oncocytic variants are representing approximatively 10% of ACT (28). The diagnosis of malignancy and the prognostic assessment for these tumors are particularly challenging. A specific malignancy score referred to as the Lin-Weiss-Bisceglia score has been specifically developed for oncocytic ACT (4, 29). When malignancy is confirmed, several studies suggest that oncocytic ACC is associated with better prognosis than conventional ACC (5, 29). However, diagnostic and prognostic criteria for oncocytic ACT have been established only in small clinical series, and these tumors have never been specifically explored in pangenomic studies. In our study, oncocytic ACT did not form any specific subgroup, suggesting that the oncocytic feature is not translating any different tumor biology. Instead, oncocytic ACT all clustered within C1B and C2 clusters. A good agreement with LinWeiss–Bisceglia score was observed, with the noticeable classification of all oncocytic ACT of uncertain malignant potential within C1B carcinomas. Of note, for non-oncocytic ACT, 6/7 ACT with Weiss score of 2 and 1/4 with Weiss score of 3 clustered in C2 adenomas. This confirms the uncertain malignant potential of non-oncocytic ACT with Weiss scores between 2 and 3.

Several studies demonstrated the importance of molecular classification for ACC prognostic stratification (13, 14, 15, 16, 18). However, transcriptome profiles and their targeted surrogates were so far limited to high-quality frozen tumor samples, precluding their use in routine practice. Moving to FFPE samples is therefore an important step. In the current study, we proposed a transcriptome-based model to classify any new ACC sample into C1A/C1B prognostic groups. The prognostic value of FFPE
transcriptome classification could be validated on an independent ACC cohort. Finally, FFPE transcriptome classification was confirmed as an independent prognostic factor in multivariable models including ENSAT stage and tumor grade.

Intratumor heterogeneity is well established in ACC, both in terms of pathology (30, 31), and somatic mutations (32, 33), while DNA methylation and chromosomal alteration profiles seem less heterogeneous (33). This study provides the first evidence of transcriptome classification in different tumor regions. (A) Hematoxylin/eosin/safran staining (40× magnification) of the two different tumor regions sampled for five patients in the training cohort, with one aggressive (region a) and one less aggressive (region b) region. The black bar represents 50 µm. Black arrows indicate mitosis. White arrows indicate atypical mitoses. (B) Description of the two different tumor regions sampled for the five patients with one aggressive (-a) and one less aggressive (-b) region. N/A: Fuhrman nuclear grade is not applicable for oncocytic cells. (C) Samples projection based on the two principle components (Dim1 and Dim2) of the PCA performed based on the 2000 most variable genes in the 95 unique patients of the training cohort. Samples from the training cohort are presented as faint circles colored by transcriptome class. Samples from two different tumor regions for eight patients are presented as full triangles colored by patient.
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homogeneity in different tumor regions. Remarkably, for five patients presenting one aggressive and one less aggressive tumor region, with mainly 'benign' features on pathology, transcriptome classification was “malignant” for both components. This is in agreement with current pathological practice, which takes into account the most aggressive tumor part for the diagnosis. The impact of sampling regions on transcriptome classification seems limited. This finding is important for using transcriptome information in clinical decisions.

This study presents some limitations. First, the sample size is limited. Further studies on larger cohorts are needed to confirm our results, especially regarding the transcriptome of oncocytic tumors and intratumor heterogeneity. However, ACC are rare, and this study included 89 ACC samples from two distinct centers. In addition, the prognostic value of FFPE transcriptome could be validated on a cohort independent from the training cohort used to establish the transcriptome classification. Secondly, a gold standard is missing for malignancy diagnosis of ACT. Indeed the current pathological scoring systems (Weiss and Lin–Weiss–Bisceglia) were established by comparing recurrent (metastazing) to non-recurrent ACT in small series of patients (2, 3, 4, 29). However, whole transcriptome classification is unsupervised and respects the data structure reflecting tumor biology.

Starting from FFPE samples, we used a 3’-end RNA-sequencing transcriptome technology. This technology is robust, cheaper and simpler to run compared to full-length RNA-sequencing (19). Other gene expression techniques starting from FFPE have been reported in ACC (34, 35, 36). However, these techniques have not emerged yet as optimal for prognostic determination in ACC. The 3’ RNA-sequencing recently emerged as an opportunity to study transcriptome signatures from FFPE material, as reported for several tumor types (37, 38, 39). This technique was applied to samples from operated patients. Whether it would be applicable to smaller amounts of tissue material,

Figure 3
FFPE transcriptome classification and prognosis in the ACC validation cohort. (A) Samples projection based on the two principle components (Dim1 and Dim2) of the PCA performed based on the 2000 most variable genes in the 95 unique patients of the training cohort. Samples from the training cohort are presented as faint circles colored by transcriptome class. Samples from validation cohort are presented as full triangles colored by transcriptome class. (B) Disease-free survival according to the predicted transcriptome class in stage I–III ACC patients of the validation cohort. (C) Overall survival according to the predicted transcriptome class in stage I–IV ACC patients of the validation cohort.

Table 2
Disease-free survival analyses in stage I-III ACC patients. P-values < 0.05 are indicated in bold.

|                      | Univariate analysis | Multivariable analysis |
|----------------------|---------------------|------------------------|
|                      | HR                  | 95% CI                 | P-value | HR                  | 95% CI                 | P-value |
| Age (per year increase) | 1.02 | 1.00–1.04 | 0.11 | 3.38 | 1.61–7.12 | < 0.001 |
| Sex (M vs F) | 0.92 | 0.45–1.86 | 0.81 | 1.36 | 0.61–3.00 | 0.45 |
| Cortisol secretion (Yes vs no) | 1.81 | 0.86–3.80 | 0.12 | 1.76 | 0.52–5.96 | 0.36 |
| ENSAT stage (III vs I–II) | 3.31 | 1.72–6.38 | 0.0003 | 3.12 | 1.02–9.57 | 0.046 |
| Resection status (R1–R2–RX vs R0) | 4.35 | 2.12–8.93 | < 10⁻⁴ | 3.39 | 1.09–10.55 | 0.035 |
| Ki-67 proliferation index | 2.85 | 0.92–8.88 | 0.07 | 1.76 | 0.52–5.96 | 0.36 |
| 10–19% vs <10% | 7.63 | 3.08–18.93 | < 10⁻⁴ | 3.12 | 1.02–9.57 | 0.046 |
| ≥20% vs <10% | 6.98 | 2.89–16.87 | < 10⁻⁴ | 3.39 | 1.09–10.55 | 0.035 |

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such as those from core biopsies, remains to be established. In this study, we based our classification of individual samples on the global transcriptome signature, a robust feature even when some individual markers are not detected because of poor RNA quality. Other advantages of FFPE samples use include an easier transfer of samples to oncogenetic departments in routine practice, the ability to optimize the tumor region to be analyzed, and the inclusion of large archive collections.

In conclusion, reliable transcriptome classification for diagnosis and prognosis of ACT can be determined from standard FFPE samples using the recent 3' RNA-sequencing technologies. This novel approach unraveled the molecular specificities of oncogenic variants. Transcriptome classification was robust in spite of intratumor heterogeneity. Finally, an accurate predictive model of transcriptome class was designed and validated for the classification of any new sample. This study provides researchers and clinicians with a convenient solution for the determination of ACT transcriptome from FFPE samples.

### Supplementary materials

This is linked to the online version of the paper at [https://doi.org/10.1530/EJE-21-1228](https://doi.org/10.1530/EJE-21-1228).

### Declaration of interest

Martin Fassnacht and Guillaume Assie are on the editorial board of EJE. They were not involved in the review or editorial process for this paper, on which they are listed as authors. The other authors have nothing to disclose.

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### Author contribution statement

J Bertherat and G Assie jointly supervised the work reported here.

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