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Authors
Winterhoff, B
Hamidi, H
Wang, C
et al.

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Molecular classification of high grade endometrioid and clear cell ovarian cancer using TCGA gene expression signatures

Boris Winterhoffa,1, Habib Hamidib,1, Chen Wangc, Kimberly R. Kallic, Brooke L. Fridleyd, Judy Deringb, Hsiao-Wang Chenb, William A. Clibye, He-Jing Wangf, Sean Dowdye, Bobbie S. Gostoute, Gary L. Keeneeye, Ellen L. Goodec, Gottfried E. Konecnyb,⁎

a Department of Obstetrics/Gynecology & Women's Health, University of Minnesota, Minneapolis, MN, United States
b Division of Hematology-Oncology, Department of Medicine, David Geffen School of Medicine, University of California Los Angeles, Los Angeles, CA, United States
c Department of Health Sciences Research, Mayo Clinic, Rochester, MN, United States
d Department of Biostatistics, University of Kansas, Kansas City, KS, United States
e Department of Gynecologic Surgery, Mayo Clinic, Rochester, MN, United States
f Department of Biostatistics, David Geffen School of Medicine, University of California Los Angeles, Los Angeles, CA, United States

HIGHLIGHTS
• TCGA gene signatures are present in high grade case of rare histology.
• Early stage high grade cases of rare histology cluster separately from advanced stages.
• TCGA signatures may help to stratify patients with rare high grade ovarian cancer.

ABSTRACT
Background. It is unclear whether the transcriptional subtypes of high grade serous ovarian cancer (HGSOC) apply to high grade clear cell (HGCCCOC) or high grade endometrioid ovarian cancer (HGOEC). We aim to delineate transcriptional profiles of HGCCCOCs and HGOECs.

Methods. We used Agilent microarrays to determine gene expression profiles of 276 well annotated ovarian cancers (OCs) including 37 HGCCCOCs and 66 HGGOECs. We excluded low grade OCs as these are known to be distinct molecular entities. We applied the prespecified TCGA and CLOVAR gene signatures using consensus non-negative matrix factorization (NMF).

Results. We confirm the presence of four TCGA transcriptional subtypes and their significant prognostic relevance (p < 0.001) across all three histological subtypes (HGSOC, HGCCCOC and HGOEC). However, we also demonstrate that 22/37 (59%) HGCCCOCs and 30/67 (45%) HGOECs form 2 additional separate clusters with distinct gene signatures. Importantly, of the HGCCCOC and HGOECs that clustered separately 62% and 65% were early stage (FIGO I/II), respectively. These finding were confirmed using the reduced CLOVAR gene set for classification where most early stage HGCCCOCs and HGOECs formed a distinct cluster of their own. When restricting the analysis to the four TCGA signatures (ssGSEA or NMF with CLOVAR genes) most early stage HGCCCOCs and HGOEC were assigned to the differentiated subtype.

Conclusions. Using transcriptional profiling the current study suggests that HGCCCOCs and HGOECs of advanced stage group together with HGSOCs. However, HGCCCOCs and HGOECs of early disease stages may have distinct transcriptional signatures similar to those seen in their low grade counterparts.

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1. Introduction
Microarray-based gene expression studies demonstrate that ovarian cancer (OC) is both a clinically diverse and molecularly heterogeneous disease, comprising subtypes with distinct gene expression patterns that are each associated with statistically significant different clinical outcomes. A gene expression analysis of high-grade serous and endometrioid OCs as part of the Australian Ovarian Cancer Study identified distinct
molecular subtypes that have been designated with neutral descriptors (C1, C2, C4, and C5) [1]. The four molecular subtypes were validated in 489 high grade serous ovarian cancer (HGSOC) cases using 1500 intrinsically variable genes for consensus non-negative matrix factorization (NMF) clustering and were termed immunoreactive, differentiated, proliferative and mesenchymal on the basis of gene expression in the clusters [2]. These four molecular subtypes have been independently validated and have been shown to be of independent prognostic relevance [3]. Using the TCGA ovarian cancer data set, Verhaak et al. recently confirmed the four molecular subtypes of high grade serous ovarian cancer (HGSOC) using a reduced subtype gene expression signature, named “Classification of Ovarian Cancer” (CLOVAR) [4]. This reduced CLOVAR gene signature is composed of a 100 genes capable of predicting the ovarian cancer subtypes [4]. Validation studies in independent data sets demonstrated that the CLOVAR signature classifies HGSOC with small error rates, making implementation using medium-throughput expression profiling platforms feasible [4].

The main objective of a molecular classification of OC into subtypes with distinct gene expression patterns is to develop robust biomarker signatures that will allow clinicians to identify women likely to benefit from a given therapy. These evolving subgroups are thought to have distinct biologic features that can translate into different therapeutic implications. Epithelial ovarian cancer is a heterogeneous disease consisting of tumors with different histology and grade. The most common OC types are the serous tumors followed by endometrioid and clear-cell cancers which represent 50%–60%, 25% and 4% of all ovarian tumors, respectively [5]. Importantly, however, the evolving molecular classification using the four main subtype signatures have almost exclusively been studied and applied to HGSOC [2,3,4]. Although some early gene expression studies have included endometrioid and clear cell ovarian cancers [6–10] these studies were limited by their small sample size and the use of early generation microarrays. Nevertheless these studies did suggest that clear cell and endometrioid ovarian cancers may be distinguished from serous ovarian cancers based on their gene expression profiles [6–10]. However, many of these early studies included well differentiated tumors (G1) known to be distinct molecular entities [11]. To date it is unclear if the evolving signatures which have been used to successfully classify HGSOC into four molecular subtypes could also be used to classify these less common epithelial ovarian cancer histologies. Although clear cell carcinomas and endometrioid carcinomas have been previously shown to be in part driven by pathways distinct from those driving progression of HGSOC we wanted to investigate whether high grade clear cell ovarian cancers (HGCCOCs) or high grade endometrioid ovarian cancers (HGEOCs) may nevertheless in part share gene signatures that have been described in HGSOCs. For instance, we thought it would be important to know if gene signatures characterizing an immunoreactive or mesenchymal subtype can also be found in HGEOCs or HGCCOCs because the evolving molecular signatures are becoming increasingly clinically relevant. In the present study we, therefore, examined the transcriptional profiles of 276 ovarian cancer cases including 37 HGCCOCs, 66 HGEOCs and 173 previously published HGSOCs using Agilent Whole Human Genome 4x44K Expression Arrays [3]. All low grade tumors were excluded from this study as they are known to represent distinct biologic entities [11]. We applied the pre-specified TCGA gene expression signatures and the reduced CLOVAR gene signatures to this cohort of 276 well annotated OCs from Mayo Clinic. Moreover, we also performed single sample gene set enrichment analysis (ssGSEA) which calculates separate enrichment scores for each sample and allows the assignment to the nearest TCGA subgroup.

2. Materials and methods

2.1. Patient cohort

Fresh frozen tumors were collected from a series of 276 consecutive women with high grade serous, clear cell and endometrioid ovarian, primary peritoneal or fallopian tube cancer who underwent surgery by a gynecologic oncologist at Mayo Clinic between 1994 and 2005. All patients signed an Institutional Review Board approved consent for bio-banking, clinical data extraction, and molecular analysis. Clinical data were abstracted from medical records and tumor registry. Thirteen patients (7.5%) were included in the TCGA study.

2.2. Sample processing and gene expression profiling

Samples were collected during surgery, snap frozen within 30 min, and stored at —80 °C until RNA extraction. Samples were reviewed by a pathologist specialized in gynecologic oncology (G.K.) and selected to have >70% tumor cell content. RNA was isolated using RNaseq (Qiagen Inc., Valencia, CA) and quantified using a Nanodrop Spectrophotometer (Agilent Technologies, Santa Clara, CA). Gene expression profiles were established using Agilent Whole Human Genome 4x44K Expression Arrays. Total RNA (750 ng) with RNA Integrity Number > 8.0 was labeled with cyanine 5-CTP or cyanine 3-CTP using the Low RNA Input Fluorescent Linear Amplification Kit (Agilent Technologies), purified on RNeasy Mini columns (Qiagen Inc.), and hybridized to expression arrays (using a mixed reference containing equal amounts of each of 106 ovarian tumor samples). Slides were scanned using the Agilent 2565BA Scanner and data were exported by the Agilent Feature Extraction Software (version 7.5.1) into Rosetta Resolver (Rosetta Inpharmatics LLC, Cambridge, MA). Log ratios of signal from individual tumor to signal from the reference mix were used for analysis. The data have been deposited in NCBI’s Gene Expression Omnibus and are accessible through GEO Series accession number: GSE73614.

2.3. Non-negative matrix factorization (NMF) and class prediction

Molecular classification was determined blinded to demographic and clinical information. To identify genes associated with TCGA subtypes, we analyzed expression and molecular subtype data of TCGA cases (https://tcga-data.nci.nih.gov/docs/publications/ov_2011/). Next, we mapped these signature genes to corresponding Agilent probe-set IDs. We selected 1844 probes matching the TCGA signature gene set. Subclasses were computed by reducing the dimensionality of the expression data from thousands of genes to a few metagenes using a consensus NMF clustering method [12]. This method computes multiple k-factor factorization decompositions of the expression matrix and evaluates the stability of the solutions using a cophenetic coefficient. The same analysis was repeated using 161 probes representing 100 genes that represented the CLOVAR subtype signature derived by Verhaak and colleagues [4].

2.4. Gene set enrichment analysis

Gene set activation scores for each of the subtype expression signatures were generated using single sample Gene Set Enrichment Analysis (ssGSEA) [13] using Bioconductor package GSVA downloadable from http://www.bioconductor.org [14]. Raw enrichment scores were expressed as relative z-scores. Subtype assignment of each tumor sample was determined using a z-score cutoff 0.6.

2.5. Statistical analysis of molecular subtypes and patient outcome

Subgroup assignments were compared by use of the chi-square test. Overall survival is depicted according to the method of Kaplan and Meier, and the curves were compared with use of the log rank test. All statistical tests were two-sided.

3. Results

We used Agilent microarrays to determine gene expression profiles of 276 well annotated OCs including 37 HGCCOCs and 66 HGEOCs and
173 HGSOCs. Patient and disease characteristics for the entire cohort and the histologic subtypes are shown in Table 1. Median patient age was 63 years (range 24–89). In the overall cohort of 276 patients 56 (21%) of the patients had early stage (FIGO I or II) disease, while 220 (79%) were diagnosed with advanced stage disease (FIGO III or IV). In the cohort of HGEOC, HGCCOC, and HGSOC 28/66 (33%), 20/37 (54%) and 8/173 (5%) of the patients had early stage disease, respectively. All tissue specimens were centrally reviewed by a pathologist specialized in gynecologic oncology (G.K.) for confirmation of tissue morphology and histologic grade. All samples were of high grade (12% G2 and 88% G3) (Table 1). In the cohort of HGEOC, HGCCOC, and HGSOC 26/66 (39%), 3/37 (8%) and 2/173 (2%) of the patients had grade 2 disease, respectively (Table 1).

A gene set representing the pre-specified TCGA signatures was used to classify 276 arrays representing HGSOC (n = 173) and HGCCOC (n = 37) and HGEOC (n = 66). We confirmed the presence of the immunoreactive, differentiated, proliferative and mesenchymal transcriptional subtypes and their prognostic relevance (p < 0.001) across all three histological subtypes (HGSOC, HGCCOC and HGEOCs) (Fig. 1). However, we also demonstrated that 22/37 (59%) HGCCOCs and 30/67 (45%) HGEOS formed two additional separate clusters with distinct gene signatures (Supplemental Tables 1 and 2). Importantly, of the HGCCOC and HGEOCs that clustered separately 62% and 65% were of early FIGO stage (FIGO I/II), respectively (Table 2). Analysis of grade distribution across the transcriptional subtypes also demonstrated that of the HGCCOC and HGEOS that clustered separately 21% and 50% were grade 2 tumors, respectively (Table 2).

Survival was significantly better for patients with tumors that demonstrated an endometrioid-like or a clear cell-like signature when compared to those patients with tumors that displayed one of the established TCGA signatures (Fig. 1). However, the good median survival in the endometrioid-like and clear cell-like cluster of 133 and 160 months, respectively, was clearly driven by early stage disease. When performing a survival analysis separately for stage III/IV patients the median survival of the endometrioid-like and clear cell-like cluster was reduced from 133 to 90 months and from 160 to 23 months, respectively. The survival analysis that was performed separately for stage III/IV patients still showed improved outcome for the endometrioid-like and clear cell-like cluster but, however, the survival for the clear cell-like cluster was comparable with the remaining poor outcome groups of HGSO when limiting the analysis to stage III/IV patients. These results underscore the notion that improved survival of the endometrioid-like or clear cell-like cluster was due to the increased rate of stage I/II diagnosis.

These findings were confirmed using the reduced CLOVAR gene set for classification into 5 subgroups where most early stage HGCCOCs and HGEOCs formed a distinct combined cluster of their own (Fig. 2). We tested the hypothesis whether gene expression based molecular subtyping of epithelial ovarian cancer developed for high grade serous histology could be applied to high grade endometrioid and clear cell histologies. Using a gene set representing the 1500 genes described by the TCGA that defined the immunoreactive, differentiated, proliferative and mesenchymal subtypes, consensus NMF revealed six distinct expression subtypes. In addition to the four TCGA subtypes we describe an endometrioid-like and a clear cell-like expression subtype when using the 1500 most variable genes described by the TCGA. When using the smaller CLOVAR gene set for consensus NMF it is possible that the smaller CLOVAR gene set for consensus NMF it is possible that more subtle difference between HGCCOC and HGEOC, which may require a bigger number of genes to be detected, may not be captured by the reduced gene list. Nevertheless, the CLOVAR signature did group many HGCCOCs and HGCCOCs into one new distinct group that was enriched for cases with early stage disease. Using transcriptional profiling the current study suggests that many HGCCOCs and HGCCOCs of advanced stage group together with HGCCOCs. However, HGCCOCs and HGCCOCs of early disease stages may have distinct transcriptional signatures more similar to those seen in their low grade counterparts. Finding from previous studies support this hypothesis [10,15–17]. Wu and colleagues identified two distinctive subgroups of endometrioid carcinoma, based on variance in their global gene expression patterns.

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Patient and disease characteristics for 276 high grade epithelial ovarian cancer patients.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HGEOC (n = 66)</td>
</tr>
<tr>
<td>Age at diagnosis</td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>58</td>
</tr>
<tr>
<td>Range</td>
<td>38–86</td>
</tr>
<tr>
<td>Grade</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>26 (39%)</td>
</tr>
<tr>
<td>3</td>
<td>40 (61%)</td>
</tr>
<tr>
<td>FIGO stage</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>19 (29%)</td>
</tr>
<tr>
<td>2</td>
<td>9 (14%)</td>
</tr>
<tr>
<td>3</td>
<td>31 (47%)</td>
</tr>
<tr>
<td>4</td>
<td>7 (10%)</td>
</tr>
<tr>
<td>Debulking status</td>
<td></td>
</tr>
<tr>
<td>Optimal</td>
<td>58 (88%)</td>
</tr>
<tr>
<td>Suboptimal</td>
<td>5 (7%)</td>
</tr>
<tr>
<td>Unknown</td>
<td>3 (5%)</td>
</tr>
</tbody>
</table>

Table 2

Survival was significantly better for patients with tumors that demonstrated an endometrioid-like or a clear cell-like signature when compared to those patients with tumors that displayed one of the established TCGA signatures (Fig. 1). However, the good median survival in the endometrioid-like and clear cell-like cluster of 133 and 160 months, respectively, was clearly driven by early stage disease. When performing a survival analysis separately for stage III/IV patients the median survival of the endometrioid-like and clear cell-like cluster was reduced from 133 to 90 months and from 160 to 23 months, respectively. The survival analysis that was performed separately for stage III/IV patients still showed improved outcome for the endometrioid-like and clear cell-like cluster, however, the survival for the clear cell-like cluster was comparable with the remaining poor outcome groups of HGSO when limiting the analysis to stage III/IV patients. These results underscore the notion that improved survival of the endometrioid-like or clear cell-like cluster was due to the increased rate of stage I/II diagnosis.

4. Discussion

We tested the hypothesis whether gene expression based molecular subtyping of epithelial ovarian cancer developed for high grade serous histology could be applied to high grade endometrioid and clear cell histologies. Using a gene set representing the 1500 genes described by the TCGA that defined the immunoreactive, differentiated, proliferative and mesenchymal transcriptional subtypes and their prognostic relevance across all three histological subtypes (HGSOC, HGCCOC and HGEOCs) (Fig. 2, p < 0.001). In addition, 47/103 (46%) of HGCCOCs and HGEOCs form an additional fifth separate cluster. Importantly, of these HGCCOC and HGEOCs that clustered separately 70% were of early stage (FIGO I/II), and 43% were grade 2, respectively (Table 2). Survival was significantly better for patients with tumors that express the endometrioid-like/clear cell-like signature when compared to those patients with tumors that express one of the TCGA signatures but when the analysis was restricted to stage III/IV patients it again became apparent that this was due to the increased rate of stage I/II diagnosis.

Furthermore, using the 100 CLOVAR genes described by Verhaak and colleagues for consensus NMF we were able to demonstrate stable clustering of all cases into four transcriptional subtypes (Fig. 3 and Supplementary Fig. 1). Consensus matrices and sample correlation matrices are shown for k = 2 to k = 8 and depict stable clustering into four groups (Cophrenic coef. 0.994) (Supplementary Fig. 1). Overall survival analysis confirmed the prognostic significance of these four TCGA transcriptional subtypes (Fig. 3, p < 0.001). Patients whose tumors expressed the immunoreactive signature demonstrated the best overall survival. This appears to contrast with earlier reports where patients whose tumors expressed the immunoreactive signature demonstrated the best overall survival [3,4]. However, this may be explained by the fact that in the present study 56/103 HGCCOC and HGEOC cases were assigned to the differentiated subtype when using the four group assignment which is expected to positively impact outcome of the differentiated subgroup (Table 2).

Single-sample Gene Set Enrichment Analysis (ssGSEA) calculates separate enrichment scores for each pairing of a sample and gene set. Each ssGSEA enrichment score represents the degree to which the genes in a particular gene set are coordinately up- or down-regulated within a sample and it allows the assignment of an individual sample to the nearest or closest transcriptional subtype. In the present analysis we used the CLOVAR gene set for ssGSEA. Consistent with the stable clustering into four groups using NMF classification, ssGSEA allowed the assignment of each HGCCOC or HGEOC to one of the four known TCGA subgroups which showed a high concordance with the consensus NMF clustering into four groups (p < 0.001, data not shown). Similar to the results obtained by consensus NMF, ssGSEA assigned 47/103 HGCCOC and HGEOC cases to the differentiated subtype.
One of these subgroups was highly similar to serous carcinoma and tended to be of higher grade. Genetic annotation also revealed that p53 mutations were common among those endometrioid carcinomas with a serous-like gene expression profile [15]. Moreover, deregulated β-catenin signaling and defects in the PI3K–PTEN pathway were shown to be typical among those endometrioid carcinomas that did not share gene expression homology to serous carcinoma [15]. In a further analysis, WT1 gene expression was associated with those endometrioid ovarian cancers with a serous-like expression profile [16]. Further studies will be necessary to understand whether these molecular differences can also be identified in the current cohort of HGEOCs. These molecular features may ultimately help us to better delineate serous from endometrioid histologies as histopathological diagnosis of high-grade endometrioid and serous cancer of the ovary is poorly reproducible under the current morphology based classification system, especially for anaplastic, high-grade tumors.

Previous studies suggest that clear cell cancer of the ovary is a unique ovarian cancer subtype [18]. Recent work has shown that a subset of clear cell cancers evolve from endometriosis [19]. Molecular alterations within clear cell cancers include unique cytokine expression patterns and c-met amplification as well as mutations in ARID1A and the PI3K/PTEN pathway [20–22]. Clear cell-specific clinical trials based upon these biologic discoveries have been designed and are presently active [23]. Furthermore, a recent study suggests that clear cell ovarian cancers have distinct molecular characteristics especially with regards to IL6-STAT3-HIF signaling and possible susceptibility to angiogenesis inhibition when compared to high grade serous and endometrioid histologies [10]. In our data set, early stage clear cell cancer of high grade had a distinct gene expression profile, confirming earlier reports on the distinct features of clear cell ovarian cancer. However, we were also able to show that many HGCCOCs of advanced stage grouped together with HGSOCs. This may in part be explained by the fact that.

![Fig. 1. Unsupervised clustering (non-negative matrix factorization) using 1500 TCGA genes depicting six clusters; overall survival according to transcriptional subtypes.]

**Table 2**
Cross tabulation assignment of high grade clear cell ovarian cancers and endometrioid ovarian cancers to the molecular subtypes (TCGA 6 clusters; CLOVAR 5 and 4 clusters) according to their FIGO stage (FIGO I/I versus III/IV) and grade (2 versus 3).

<table>
<thead>
<tr>
<th>FIGO stage</th>
<th>Grade</th>
<th>P value</th>
<th>P value</th>
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</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>I/II</td>
<td>III/IV</td>
</tr>
<tr>
<td>N</td>
<td>103</td>
<td>48</td>
<td>55</td>
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<table>
<thead>
<tr>
<th>TCGA 6*</th>
<th></th>
<th></th>
<th></th>
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<th></th>
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</thead>
<tbody>
<tr>
<td>Immunoreactive</td>
<td>12 (100%)</td>
<td>4 (33%)</td>
<td>8 (67%)</td>
<td>0.001</td>
<td>2 (17%)</td>
<td>10 (83%)</td>
</tr>
<tr>
<td>Differentiated</td>
<td>15 (100%)</td>
<td>1 (7%)</td>
<td>14 (93%)</td>
<td></td>
<td>2 (13%)</td>
<td>13 (87%)</td>
</tr>
<tr>
<td>Proliferative</td>
<td>12 (100%)</td>
<td>3 (25%)</td>
<td>9 (75%)</td>
<td></td>
<td>3 (25%)</td>
<td>9 (75%)</td>
</tr>
<tr>
<td>Mesenchymal</td>
<td>6 (100%)</td>
<td>3 (50%)</td>
<td>3 (50%)</td>
<td></td>
<td>0</td>
<td>6 (100%)</td>
</tr>
<tr>
<td>Clear cell-like</td>
<td>24 (100%)</td>
<td>15 (62%)</td>
<td>9 (38%)</td>
<td>5 (21%)</td>
<td>19 (79%)</td>
<td></td>
</tr>
<tr>
<td>Endometrioid-like</td>
<td>34 (100%)</td>
<td>22 (65%)</td>
<td>12 (35%)</td>
<td>17 (50%)</td>
<td>17 (50%)</td>
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<table>
<thead>
<tr>
<th>CLOVAR 5#</th>
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<th></th>
<th></th>
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<tbody>
<tr>
<td>Immunoreactive</td>
<td>15 (100%)</td>
<td>3 (20%)</td>
<td>12 (80%)</td>
<td>0.001</td>
<td>1 (7%)</td>
<td>14 (93%)</td>
</tr>
<tr>
<td>Differentiated</td>
<td>19 (100%)</td>
<td>6 (32%)</td>
<td>13 (68%)</td>
<td></td>
<td>4 (21%)</td>
<td>15 (79%)</td>
</tr>
<tr>
<td>Proliferative</td>
<td>15 (100%)</td>
<td>4 (27%)</td>
<td>11 (73%)</td>
<td></td>
<td>4 (27%)</td>
<td>11 (73%)</td>
</tr>
<tr>
<td>Mesenchymal</td>
<td>7 (100%)</td>
<td>2 (29%)</td>
<td>5 (71%)</td>
<td></td>
<td>0</td>
<td>7 (100%)</td>
</tr>
<tr>
<td>Clear cell-like/endometrioid-like</td>
<td>47 (100%)</td>
<td>33 (70%)</td>
<td>14 (30%)</td>
<td>20 (43%)</td>
<td>27 (57%)</td>
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</tbody>
</table>

<table>
<thead>
<tr>
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<th></th>
<th></th>
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<th></th>
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</thead>
<tbody>
<tr>
<td>Immunoreactive</td>
<td>18 (100%)</td>
<td>5 (28%)</td>
<td>13 (72%)</td>
<td>0.037</td>
<td>1 (6%)</td>
<td>17 (94%)</td>
</tr>
<tr>
<td>Differentiated</td>
<td>56 (100%)</td>
<td>33 (59%)</td>
<td>23 (41%)</td>
<td></td>
<td>23 (41%)</td>
<td>33 (59%)</td>
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<tr>
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<td>8 (40%)</td>
<td>12 (60%)</td>
<td></td>
<td>5 (25%)</td>
<td>15 (75%)</td>
</tr>
<tr>
<td>Mesenchymal</td>
<td>9 (100%)</td>
<td>2 (22%)</td>
<td>7 (78%)</td>
<td></td>
<td>0</td>
<td>9 (100%)</td>
</tr>
</tbody>
</table>

*Assignment using unsupervised clustering with the 1500 most differentially expressed genes described by the TCGA.
#Assignment using unsupervised clustering with the reduced gene set CLOVAR genes described by Verhaak et al.
the TCGA expression signatures may reflect disease processes that are more critical for advanced ovarian cancer than for early disease stages. While it is evident that clear cell carcinomas are in part driven by pathways distinct from those driving progression of HGSOc, they may, nevertheless, in part share gene signatures that have been described in HGSOc. As the evolving molecular signatures in HGSOc are becoming increasingly clinically relevant it is important to understand whether therapeutically relevant gene signatures like the immunoreactive or mesenchymal signature can also be found in HGCCOCs. As the evolving molecular signatures in HGSOc are becoming increasingly clinically relevant it is important to understand whether therapeutically relevant gene signatures like the immunoreactive or mesenchymal signature can also be found in HGCCOCs.

Consensus NMF and ssGSEA using the reduced CLOVAR gene set both likewise demonstrated stable clustering of all OC cases into four subgroups. Importantly, we were able to show that HGCCOC and HGEOC cases of early stage and G2 tended to fall into the differentiated expression subgroup and that HGCCOCs and HGEOCs of advanced stage more likely group together with the remaining HGSOcs transcriptional subtypes. Importantly, however, before any of the proposed classifications can be converted to clinical use, further clinical validation of their predictive importance will be key in moving either of them forward. Associations between subtype signatures and treatment responses will help to clarify which signatures can be deemed to be the most appropriate to help classify OC. Only these studies will allow us to ultimately determine which subtype signatures are most appropriate to select patients for any given therapy. Possibly these evolving ovarian cancer transcriptional signatures will be used as predictive signatures similar to how gene expression based classifiers are currently being used to guide treatment decisions in breast cancer [24]. Results from a recent retrospective study support this notion. Using NMF to assign the gene expression profiles of 359 formalin fixed paraffin embedded tumor samples from patients treated in a phase III frontline ovarian cancer trial, which assessed the efficacy of bevacizumab, into the four molecular subtypes demonstrated a preferential treatment effect of bevacizumab in the proliferative and mesenchymal subtypes [25]. These preliminary data underscore the potential clinical utility of transcriptional signatures across all advanced ovarian cancer types including HGCCOCs and HGEOCs. Our study, however, is not without limitations as neither additional genomic nor immunohistochemical analyses have been performed to further delineate the biologic underpinnings of the distinct transcriptional signatures seen between advanced and early stage HGCCOCs and HGEOCs. Nevertheless, a strength of our study is the fact that we were able to profile a relatively large number of rare high grade epithelial ovarian cancers which allowed us to demonstrate the existence of the immunoreactive, differentiated, proliferative and mesenchymal transcriptional subtypes across advanced ovarian cancer cases irrespective of their specific
histology. Importantly, however, rare high grade epithelial ovarian cancers of early disease stages may have distinct transcriptional signatures similar to those seen in their low grade counterparts. These findings could aid in the implementation of medium-throughput expression profiling platforms to guide treatment decisions in advanced epithelial ovarian cancer and may support their use across different histologies.

Conflict of interest

The authors declare that there are no conflicts of interest.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.jygyno.2016.02.023.

References