Review Article

Prognostic Significance of Vascular Endothelial Growth Factor Serum Determination in Women with Ovarian Cancer

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1. Introduction

Ovarian cancer is the most frequent cause of death from gynaecological cancer and the fourth most frequent cause of cancer-related death in women in Europe and the United States [1]. It has the highest fatality-to-case ratio of all gynaecological malignancies, mainly due to the fact that it is characterized by early widespread metastasis and high-grade malignancy at diagnosis. The five-year survival proportion is about 80–90% for patients with stage I disease and only 15–20% for patients with stage III or IV disease. Although survival has improved with the use of maximal cytoreduction surgery along with platinum- and taxane-based chemotherapy, nearly 80% of ovarian cancers relapse and patients inevitably succumb to the development of chemotherapy-resistant disease [2].

Clinicopathological features known to be prognostic variables for ovarian cancer are surgical stage (FIGO stage), histological grade, lymph node involvement, residual tumour size after cytoreductive surgery, histological subtype, ascites, and age. According to the three-year analysis of the FIGO Annual Report on the Results of Treatment in Gynaecological Cancer, stage, grade, and residual tumour size have the greatest prognostic value [3]. However, these
factors provide an insufficient picture of the biology of ovarian cancer and they are frequently interrelated. The identification of new serological biological markers that predict the outcome of the disease would be extremely useful for developing individually tailored and possibly more effective treatments. Serum analysis is a noninvasive technique feasible in cases where no tissue is available and it can also be performed during followup.

It is well established that angiogenesis, the formation of new blood vessels, is necessary for the growth and metastatic spread of solid tumours [4–7]. A high degree of tumour angiogenesis has been shown to correlate with poor survival in women with ovarian cancer [7–10]. Vascular endothelial growth factor (VEGF) plays an essential role in angiogenesis in many tumour types [11–15]. It is a heparin-binding dimeric glycoprotein involved in angiogenic, mitotic, and microvascular permeability-inducing activities, leading to extravasation of plasma proteins and proangiogenic stromal changes [16].

Several studies have found VEGF levels to be significantly higher in the tissues and biological fluids of women with ovarian cancer compared with healthy controls [17–20], whereas its association with tumour progression and/or patient survival is still controversial. We performed a review of the literature to elucidate the prognostic role of serum VEGF (sVEGF) levels in ovarian cancer, both alone and in comparison with established clinicopathological factors.

2. Methods

Eligible studies in English and Italian were identified in MEDLINE (PubMed version) from VEGF discovery to October 2011 using the terms VEGF, “vascular endothelial growth factor” and synonyms, “ovarian cancer,” “ovary cancer,” and synonyms. We searched studies that used these terms in title and abstract and that was indexed by bibliographic database with the Mesh term “ovarian neoplasms.” We searched the database using these terms separately and also in combination.

Relevant papers were independently selected by two of the reviewers (E. Bandiera and R. Franceschini) based on the following inclusion criteria: studies that evaluated (i) sVEGF levels before any surgical and chemotherapeutic treatment; (ii) the association of sVEGF levels with the established clinicopathological prognostic factors (FIGO stage, tumour grade, residual tumour size, lymph node involvement, histological type, ascites, age); (iii) the value of sVEGF levels in predicting patients’ outcomes (overall survival (OS), disease-free survival (DFS), progression free survival (PFS)). Any disagreement in the inclusion of one study between two reviewers was solved by discussion.

2.1. Data Extraction. Two of the reviewers (E. Bandiera and R. Franceschini) independently reviewed each study and abstracted data on first author, country of study, study characteristics (study design, followup duration, therapy), clinical and pathological variables, and study outcomes.

3. Results

3.1. Included Studies. Our search strategy identified 758 journal abstracts. From these, we retrieved for further evaluation 15 full-text articles focused on the relationship between circulating preoperative VEGF and prognosis in ovarian cancer. Of these 15 articles, nine [21–29] studies met the inclusion criteria and were used for this review.

Hefler et al. [28] gathered together some cases from trials by Tempfer et al. [21], Gadducci et al. [22], Chen et al. [23], and Cooper et al. [25]. Because Hefler et al. [28] added a series of new patients, we reported these five [21–23, 25, 28] studies independently.

3.2. Excluded Studies. Six studies were excluded. The paper by Manenti et al. [30] was excluded because these authors analysed plasma VEGF levels. Since VEGF is secreted also by platelets, Manenti et al.’s study was unsuitable for comparison with studies focusing on sVEGF. Boss et al. [31], Bamias et al. [32], Yamamoto et al. [33], and Rudlowski et al. [34] were excluded because they did not directly evaluate the prognosis of patients with abnormal sVEGF levels and/or the association between sVEGF levels and clinicopathological characteristics. Finally, Dirix et al. [35] evaluated patients with different cancer types and did not report independent results for ovarian cancer.

3.3. Characteristics of the Selected Studies. The nine selected studies were published between 1996 and 2010 and included 529 patients from seven countries. All but one [28] study were retrospective. Six of these nine studies reported details on the duration of followup (median: 42 months [21], 34 months [23], 29 months [29] or mean: 39 months [28], or length: 60 months [25], 24 months [22]). All studies analysed the association between sVEGF levels and the established prognostic variables and evaluated the association between sVEGF levels and OS. Three studies [21, 23, 27] analysed the association between sVEGF and DFS whereas only one [29] analysed the association between sVEGF and PFS.

3.4. Characteristics of Study Populations. All studies enrolled women with newly diagnosed and histopathologically confirmed ovarian cancer except the one by Cooper et al. [25], which included also a small group of women with peritoneal and fallopian tube malignancies (Table 1). The mean (or median) ages of the studied women ranged from 52.5 to 64 years. Most cancers (>95%) were epithelial and the predominant histological type was serous carcinoma. Most patients were diagnosed with poorly differentiated ovarian cancer, advanced FIGO stage, and ascites. With the exception of Gadducci et al. [22], who followed only 27 patients with advanced disease receiving chemotherapy, all other studies monitored all patients enrolled.

Surgery for optimal tumour debulking included hysterectomy, bilateral salpingo-oophorectomy, omentectomy [21, 23, 28, 29], pelvic and para-aortic lymphadenectomy [21, 23, 28, 29], and appendectomy [23, 28, 29]. Five studies [22, 24–27] did not describe the surgical approach, but
| Origin and Dates       | Austria 1990–1995 | Italy 1990–1997 | Taiwan 1992–1998 | Germany 1995–2000 | Iowa 1999–2001 | China 1997–2002 | Poland 1990–2003 | Austria, Iowa, Italy, Taiwan 1996–2004 | Germany 1996–2004 |
|-----------------------|-------------------|-----------------|------------------|-------------------|----------------|-----------------|-----------------|----------------------------------------|------------------|
| Number of cases       | 60                | 53              | 56               | 41                | 101            | 50              | 86              | 314                                    | 37               |
| Age, mean or median* (range), years | 55.6 (36–71) | 59* (23–81) | 52.5* (21–88) | 62 (20–78) | NA            | NA              | 59.9            | 58.61* (26–78)                        |
| FIHG stage            |                   |                 |                  |                   |                |                 |                 |                                        |
| I                     | 7                 | 19              | 14               | 5                 | 20             | 17              | 14              | 56                                     | 1                |
| II                    | 12                | 1               | 6                | 2                 | 9              | 27              | 37              | 177                                    | 29               |
| III                   | 27                | 22              | 32               | 30                | 81             | 33              | 37              | 177                                    | 29               |
| IV                    | 14                | 11              | 4                | 4                 | NA             | NA              | 26              | 46                                     | 6                |
| Grade                 |                   |                 |                  |                   |                |                 |                 |                                        |
| G1                    | 21                | 17              | 21               | NA                | 19             | 19              | 60              | 0                                      |
| G2                    | 39                | 12              | 35               | NA                | 39             | 13              | 41              | 88                                     | 9                |
| G3                    |                   |                 |                  |                   |                |                 | 30              | 26                                     | 150              |
| Epithelial ovarian cancer |               |                 |                  |                   |                |                 |                 |                                        |
| Serous                | 28                | 33              | 30               | 32                | 15             | 49              | 166             | 31                                     |
| Mucinous              | 23                | 6               | 4                | 2                 | 9              | 4               | 41              | 0                                      |
| Undifferentiated      |                   |                 |                  |                   |                |                 |                 |                                        |
| Endometrioid          | 3                 | 6               | 8                | 0                 | 9              | 4               | 41              | 0                                      |
| Clear cell            | 1                 | 1               | 0                | 0                 | 0              | 0               | 9               | 1                                      |
| Others                | 1                 | 1               | 0                | 0                 | 0              | 0               | 44              | 0                                      |
| Nonepithelial ovarian cancer |         |                 |                  |                   |                |                 |                 |                                        |
| >500 mL/presence*     | NA                | 21**            | NA               | NA                | 67**           | 22              | 64**            | NA                                     | 21               |
| <500 mL/absence**     | NA                | 12**            | NA               | NA                | 34**           | 28              | 22**            | NA                                     | 15               |
| Ascites               |                   |                 |                  |                   |                |                 |                 |                                        |
| Residual disease      |                   |                 |                  |                   |                |                 |                 |                                        |
| <2 cm                 | 38                | 9               | 40               | 11                | 44             | 32              | 164***          | 22                                     |
| >2 cm                 | 22                | 24              | 16               | 30                | 6              | 54              | 64***           | 14                                     |
| Lymph node involvement|                   |                 |                  |                   |                |                 |                 |                                        |
| yes                   | 23                | NA              | NA               | NA                | 12             | NA              | 38              | 10                                     |
| no                    | 37                | NA              | NA               | NA                | 38             | NA              | 63              | 11                                     |

1: Cooper study contains a small group of peritoneal and fallopian tube malignant cancers; NA: not available data; *: Median values; **: Numbers of patients with presence/absence of ascites; ***: Numbers of patients with residual disease < or >1 cm.
according to the reported data (residual tumour size [22, 24–27], omental metastasis [26], lymph node involvement [26]), we may presume that maximal cytoreductive surgery was performed.

Surgery was followed by chemotherapy consisting of platinum analogues alone [21–24, 28] or in combination with taxane [25, 29]. Early stages of disease were treated according to the standards established by the respective institutions: patients with stage IA-IB [21], I-II [22], IA [25], and IA-IB excluding clear cell histology [28] did not receive any chemotherapy or were treated like patients with advanced disease [23, 24, 29]. Although postoperative chemotherapy is the accepted standard treatment, two studies [26, 27] omitted any information about it.

3.5. sVEGF Assay. The sVEGF assay method was similar across studies. Venous blood was taken preoperatively from all patients. All sera were separated and stored at ≤20°C. Seven studies [21–25, 27, 28] used the same Quantikine sandwich ELISA kit (R&D Systems Minneapolis, USA). Li et al.’s study used a home-made indirect ELISA kit, whereas Mahner et al.’s study used VEGF-165 ELISA KIT (Siemens Healthcare Diagnostic, Tarrytown, USA).

Data on the precision of sVEGF assays were reported in three studies [21, 24, 28], and in all studies, the intra/inter-assay coefficient of variation was <10%. Median values of sVEGF reported by authors were: 466 pg/mL [21], 229 pg/mL [22], 458 pg/mL [23], 440 pg/mL [24], 379 pg/mL [25], 387 pg/mL [27], 407 pg/mL [28], and 171 pg/mL [29]. Li et al. [26] showed a mean value of 765 pg/mL.

3.6. Relationship between sVEGF Levels and the Other Prognostic Factors. The association between sVEGF concentrations and FIGO stage, tumour grade, residual tumour size, lymph node involvement, histological type, ascites, and age was analysed by 89%, 89%, 89%, 44%, 67%, 56%, and 78% of studies respectively (Table 2).

When the median (or mean) of sVEGF values was evaluated in relation to clinicopathological features, a statistically significant association between the level of sVEGF and FIGO stage, tumour grade, residual tumour size, lymph node involvement, and presence of ascites was found in at least one study. By contrast, no statistically significant association was found between sVEGF levels and histological type or age.

Tempfer et al. [21], Chen et al. [23], and Li et al. [26] demonstrated that elevated sVEGF levels were associated with a high malignant potential of tumours (G1 versus G2-G3 [21, 23], G1-G2 versus G3 [26]). Gadducci et al. [22], Cooper et al. [25], and Li et al. [26] reported a positive association between sVEGF concentrations and ascites volume (Gadducci et al. [22] selected patients with stages III-IV). Li et al. [26] and Hefler et al. [28] found that patients with suboptimally debulked cancer had higher sVEGF values than patient in whom tumour debulking was optimal. Finally, only the study by Li et al. [26] showed that sVEGF values were higher in patients with advanced FIGO stages and lymph node involvement.

3.7. sVEGF Evaluation. In evaluating its association with outcome variables, the levels of sVEGF were dichotomised using different cut-offs: 75th percentile [21, 23] or median [24, 29] in ovarian cancer patients, mean [26] in healthy subjects, and 95th percentile [27] in patients with benign disease. In one study, the authors used the value maximizing the hazard ratio [25]. Cut-offs ranged from 100 to 826 pg/mL. Finally, only Hefler and coworkers [28] considered sVEGF as a continuous variable.

3.8. Statistical Analyses. In the univariate analyses, seven [21, 23, 24, 26–29] studies used the Kaplan-Meier product-limit method to estimate how sVEGF and other clinicopathological variables might predict OS and DFS. Gadducci et al. [22] and Cooper et al. [25] did not explicitly report the method of univariate analysis. In the multivariate analyses, all studies claim to have used the Cox proportional hazards regression model to assess the independent role of different, simultaneously evaluated prognostic factors in determining outcomes. Estimates are reported in terms of relative risk (RR) and hazard ratio (HR). The results of univariate and multivariate analyses were considered statistically significant when the P values were <0.05.

3.9. Univariate and Multivariate OS Analysis. Univariate and multivariate analyses for survival were reported in Table 3. All studies analysed the association between sVEGF levels and OS. With the exception of Gadducci et al. [22] and Mahner et al. [29], all authors found that elevated sVEGF was associated with shorter OS. Moreover, five [21, 23, 25, 27, 28] of these seven studies found sVEGF to be an independent prognostic factor.

As expected, clinicopathological features known to be prognostic variables for EOC such as FIGO stage, tumour grade, residual tumour size after cytoreductive surgery, lymph node involvement, and age have been shown as independent prognostic factors in at least one study. Notably, sVEGF, in comparison with others prognostic variables, was reported as independent prognostic factors by the majority of studies.

Chen et al. [23], Li et al. [26], and Hefler et al. [28] chose subgroups of patients for further analyses. Chen et al. [23] selected a subset of 40 patients with residual tumour size less than 2 cm. Univariate analysis, performed only for sVEGF, showed that elevated sVEGF was associated with shorter OS. Multivariate analysis identified sVEGF, FIGO stage, and grade as independent prognostic factors.

Li et al. [26] demonstrated that there was no significant difference in cumulative survival probability between stage I/II patients with high values of sVEGF and stage I/II patients with low levels of sVEGF. By contrast, the cumulative survival probability of stage III/IV patients with high values of sVEGF was lower than that of stage III/IV patients with low levels of sVEGF.

A planned subgroup analysis was performed for 56 patients with FIGO stage I in the study by Hefler et al. [28]. In univariate analysis, only sVEGF was associated with
# Table 2: Association between sVEGF and clinicopathological characteristics of patients.

| Variable               | Author, year               | No. cases | Reported statistics for VEGF | Variable type | Statistical significance of association |
|------------------------|---------------------------|-----------|------------------------------|---------------|-----------------------------------------|
| **Stage**              |                           |           |                              |               |                                         |
|                        | Tempfer et al., 1998 [21] | 60        | md                           | I/II versus III/IV | NO                                      |
|                        | Gadducci et al., 1999 [22]| 53        | md                           | I versus II and III versus IV | NO                                      |
|                        | Chen et al., 1999 [23]    | 56        | md                           | I/II versus III/IV | NO                                      |
|                        | Oehler and Caffier, 2000 [24]| 41   | m                             | categorical     | NO                                      |
|                        | Cooper et al., 2002 [25]  | 101       | md                           | I/II versus III/IV | NO                                      |
|                        | Li et al., 2004 [26]      | 50        | m                             | I/II versus III/IV | YES                                     |
|                        | Harlozińska et al., 2004 [27]| 86    | NA                           | I/II versus III/IV | NO                                      |
|                        | Hefer et al., 2006 [28]   | 314       | m                             | categorical     | NO                                      |
| **Grade**              |                           |           |                              |               |                                         |
|                        | Tempfer et al., 1998 [21] | 60        | md                           | G1 versus G2/G3 | YES                                     |
|                        | Gadducci et al., 1999 [22]| 53        | md                           | G1-G2 versus G3 | NO                                      |
|                        | Chen et al., 1999 [23]    | 56        | md                           | G1 versus G2/G3 | YES                                     |
|                        | Cooper et al., 2002 [25]  | 101       | md                           | G1-G2 versus G3 | NO                                      |
|                        | Li et al., 2004 [26]      | 50        | m                             | G1-G2 versus G3 | YES                                     |
|                        | Harlozińska et al., 2004 [27]| 86    | NA                           | G1 versus G2/G3 | NO                                      |
|                        | Hefer et al., 2006 [28]   | 314       | m                             | categorical     | NO                                      |
|                        | Mahner et al., 2010 [29]  | 37        | md                           | G2 versus G3   | NO                                      |
| **Residual tumour size (cm)** |                       |           |                              |               |                                         |
|                        | Tempfer et al., 1998 [21] | 60        | md                           | ≥2 versus <2   | NO                                      |
|                        | Gadducci et al., 1999 [22]| 53        | md                           | ≥2 versus <2   | NO                                      |
|                        | Chen et al., 1999 [23]    | 56        | md                           | ≥2 versus <2   | NO                                      |
|                        | Oehler and Caffier, 2000 [24]| 41   | m                             | ≥2 versus <2   | NO                                      |
|                        | Cooper et al., 2002 [25]  | 101       | md                           | ≥1 versus <1   | NO                                      |
|                        | Li et al., 2004 [26]      | 50        | m                             | ≥2 versus <2   | YES                                     |
|                        | Hefer et al., 2006 [28]   | 314       | m                             | ≥1 versus <1   | YES                                     |
|                        | Mahner et al., 2010 [29]  | 37        | md                           | ≥0 versus <0   | NO                                      |
| **Lymph node involvement** |                       |           |                              |               |                                         |
|                        | Tempfer et al., 1998 [21] | 60        | md                           | yes versus no  | NO                                      |
|                        | Li et al., 2004 [26]      | 50        | m                             | yes versus no  | YES                                     |
|                        | Hefer et al., 2006 [28]   | 314       | m                             | yes versus no  | NO                                      |
|                        | Mahner et al., 2010 [29]  | 37        | md                           | yes versus no  | NO                                      |
OS. In multivariate analysis, sVEGF and tumour grade were independent prognostic factors for OS.

3.10. Univariate and Multivariate DFS and PFS Analysis. Only three [21, 23, 27] of the nine included studies considered DFS as an end point. In univariate analysis, a significant association between DFS and sVEGF level was found by 2 [21, 23] out of 3 [21, 23, 27] studies. In multivariate analysis, sVEGF levels were shown to be independent prognostic factors by 2 [21, 23] out of 3 [21, 23, 27] studies. The associations between DFS and other prognostic factors were shown in Table 4.

Chen et al. [23] further evaluated DFS for 40 ovarian carcinoma patients with residual tumour size less than 2 cm, and they found that elevated sVEGF levels were significantly associated with lower DFS in univariate analysis and sVEGF levels, FIGO stage and grade were independent prognostic factors for DFS in multivariate analysis.

Finally, only Mahner et al. [29] considered PFS as an end point and he did not find a significant association between PFS and sVEGF level.

4. Discussion

The management of patients with ovarian cancer is based on established prognostic factors such as tumour stage, histological grade, and residual tumour size after cytoreductive surgery. Recently, the concept of standard chemotherapeutic treatment with platinum/taxane combination, the necessity of adjuvant chemotherapy in early stages of disease, the use of neoadjuvant chemotherapy for patients expected not to be optimally debulked at primary cytoreductive surgery and the
| Variable | Author, year | No. cases | Cut-off | Univariate analysis RR or HR, P-value | Multivariate analysis RR or HR, P-value |
|----------|-------------|-----------|---------|--------------------------------------|----------------------------------------|
|          |             |           |         |                                      |                                        |
| VEGF (pg/mL) |            |           |         |                                      |                                        |
|          | Tempfer et al., 1998 [21] | 60 | ≥826 versus <826 | RR = 2.7, P = 0.007 | RR = 2.7, P = 0.008 |
|          | Gadducci et al., 1999 [22] | 53 | NA | P = NS | NA |
|          | Chen et al., 1999 [23] | 56 | NA | P < 0.001/P = 0.006* | RR = 4.47, P ≤ 0.001; RR = 5.37*, P < 0.001* |
|          | Oehler and Caffier, 2000 [24] | 41 | ≥440 versus <440 | HR = 3.56, P = 0.026 | P = NS |
|          | Cooper et al., 2002 [25] | 101 | ≥380 versus <380 | HR = 2.13, P = 0.009 | HR = 2.08, P = 0.02 |
|          | Li et al., 2004 [26] | 50 | ≥100 versus <100 | P = 0.0085; P = 0.45; P = 0.02* | P = 0.0750 |
|          | Harlozińska et al., 2004 [27] | 86 | ≥750 versus <750 | P = 0.0169 | RR = 2.35; P = 0.02 |
|          | Hefler et al., 2006 [28] | 314 | continuous variable | P < 0.001; P < 0.001* | HR = 1.8, P = 0.03; HR = 1.1*, P = 0.001* |
|          | Mahner et al., 2010 [29] | 37 | 171 | P = 0.302 | NA |
| Stage |            |           |         |                                      |                                        |
|          | Tempfer et al., 1998 [21] | 60 | I/II versus III/IV | RR = 3.2, P = 0.007 | RR = 3.2, P = 0.001 |
|          | Chen et al., 1999 [23] | 56 | I/II versus III/IV | NA | RR = 2.08, P = 0.11; RR = 3.84*, P = 0.01* |
|          | Oehler and Caffier, 2000 [24] | 41 | I/II versus III/IV | HR = 2.24, P = 0.043 | P = NS |
|          | Cooper et al., 2002 [25] | 101 | I/II versus III/IV | HR = 10.15, P < 0.001 | HR = 9.24, P < 0.001 |
|          | Li et al., 2004 [26] | 50 | NA | NA | P = NS |
|          | Harlozińska et al., 2004 [27] | 86 | I/II versus III/IV | P = 0.0006 | RR = 4.08, P = 0.008 |
|          | Hefler et al., 2006 [28] | 314 | continuous variable | P < 0.001 | HR = 1.7, P < 0.001 |
| Grade |            |           |         |                                      |                                        |
|          | Tempfer et al., 1998 [21] | 60 | G1 versus G2/3 | RR = 1.4, P = 0.005 | RR = 1.4, P = 0.01 |
|          | Chen et al., 1999 [23] | 56 | G1 versus G2/3 | NA | RR = 2.38, P = 0.034; RR = 2.44*, P = 0.045* |
|          | Cooper et al., 2002 [25] | 101 | G1/2 versus G3 | HR = 1.36, P = 0.29 | HR = 0.86; P = 0.63 |
|          | Li et al., 2004 [26] | 50 | NA | NA | P = NS |
|          | Harlozińska et al., 2004 [27] | 86 | G1 versus G2/3 | P = 0.00079 | P = NS |
|          | Hefler et al., 2006 [28] | 314 | NA | P < 0.001; P < 0.2* | HR = 1.2, P = 0.3; HR = 3.4*, P = 0.02* |
| Variable                               | Author, year                      | No. cases | Cut-off | Univariate analysis RR or HR, P-value | Multivariate analysis RR or HR, P-value |
|----------------------------------------|-----------------------------------|-----------|---------|--------------------------------------|----------------------------------------|
| Residual tumor size (cm)               | Chen et al., 1999 [23]            | 56        | ≥2 versus <2 | NA                                    | RR = 1.34, P = 0.46                     |
|                                        | Oehler and Caffer, 2000 [24]      | 41        | 0 versus 1 + 2 | HR = 11.68, P = 0.018               | HR = 11.68, P = 0.018                   |
|                                        | Cooper et al., 2002 [25]          | 101       | 0 versus 1 + 2 | HR = 2.2, P = 0.007                | HR = 1.29, P = 0.42                     |
|                                        | Li et al., 2004 [26]              | 50        | NA       | NA                                   | P = 0.019                              |
|                                        | Harlozińska et al., 2004 [27]     | 86        | ≥2 versus <2 | P = 0.00637                          | P = NS                                 |
|                                        | Hefler et al., 2006 [28]          | 314       | ≥1 versus <1 | P < 0.001                            | HR = 1.8, P = 0.006                     |
| Lymph node involvement                | Tempfer et al., 1998 [21]         | 60        | Yes versus No | RR = 2.8, P = 0.0007                | RR = 2.8, P = 0.006                     |
|                                        | Li et al., 2004 [26]              | 50        | NA       | P = NS                               |                                        |
| Histological type                     | Chen et al., 1999 [23]            | 56        | serous/mucinous versus others        | NA                                    | RR = 0.99, P = 0.92; RR = 1.14*, P = 0.21* |
|                                        | Li et al., 2004 [26]              | 50        | NA       | P = NS                               |                                        |
|                                        | Harlozińska et al., 2004 [27]     | 86        | serous versus others                 | P = NS                                |                                        |
|                                        | Hefler et al., 2006 [28]          | 314       | serous versus others                 | P = 0.3; P = 0.6*                     | HR = 1.1, P = 0.6; HR = 1’, P = 0.9*   |
| Ascites                                | Cooper et al., 2002 [25]          | 101       | presence versus absence              | HR = 2.5, P = 0.004                   | HR = 1.28, P = 0.54                     |
| Age (years)                           | Oehler and Caffer 2000 [24]       | 41        | ≥60 versus <60                      | P = NS                                |                                          |
|                                        | Cooper et al., 2002 [25]          | 101       | NA       | HR = 1.34, P = 0.30                 | HR = 1.16, P = 0.63                     |
|                                        | Harlozińska et al., 2004 [27]     | 86        | ≥62 versus <62                      | P = 0.0478                            | RR = 2.20, P = 0.0272                   |
|                                        | Hefler et al., 2006 [28]          | 314       | continuous variable                 | P = 0.01, P = 0.8*                    | HR = 1, P = 0.9; HR = 1’, P = 0.6*     |

* Subset of 40 patients with residual tumour size ≤2 cm; ** Subset of 56 patients with stage I; § Subset of patients with stages I-II; ^ Subset of patients with stages III-IV; NA: not available data; NS: non-significant statistical analysis.
use of consolidation chemotherapy for patients at high risk of recurrence have all been questioned. The need for additional prognostic data to calibrate therapeutic tools on an individual basis in women with ovarian cancer seems obvious. In contrast to other malignancies, no therapeutic tools on an individual basis in women with ovarian carcinoma patients except those by Cooper et al. [25] and Li et al. [26], where other ovarian, peritoneal, and tubal malignancies were included. Patients underwent different chemotherapy regimens based on platinum analogues alone [21–24, 28] or in combination with taxane [25, 29]. Patients with early-stage disease were treated differently or were not treated, depending on the standards of the respective institutions. None of the studies exhaustively described followup (time, lost patients, events). Although seven [21–25, 27, 28] out of nine studies used the same sVEGF assay, one [22] of these measured significantly lower sVEGF values.

Table 4: Univariate and multivariate analyses for disease free survival.

| Variable                  | Author, year          | No. cases | Cut-off               | Univariate analyses RR, P-value | Multivariate analyses RR, P-value |
|---------------------------|-----------------------|-----------|-----------------------|---------------------------------|----------------------------------|
| VEGF (pg/mL)              | Tempfer et al., 1998  | 60        | ≥826 versus <826      | RR = 1.8, P = 0.003             | RR = 1.8, P = 0.02               |
|                           | Chen et al., 1999     | 56        | NA                    | P = 0.001, P = 0.001*           | RR = 3.34, P = 0.002; RR = 5.62*, P < 0.001* |
|                           | Harlozińska et al.,   | 314       | ≥750 versus <750      | P = NS                           | P = NS                           |
| Stage                     | Tempfer et al., 1998  | 60        | I/II versus III/IV    | RR = 1.3, P = 0.01              | RR = 1.3, P = 0.02               |
|                           | Chen et al., 1999     | 56        | I/II versus III/IV    | NA                              | RR = 2.09, P = 0.10; RR = 3.28*, P = 0.027* |
|                           | Harlozińska et al.,   | 314       | I/II versus III/IV    | P = 0.000                        | RR = 4.66, P = 0.00018           |
| Grade                     | Tempfer et al., 1998  | 60        | G1 versus G2/G3       | RR = 1.9, P = 0.03              | RR = 1.9, P = 0.04               |
|                           | Chen et al., 1999     | 56        | G1 versus G2/G3       | NA                              | RR = 2.24, P = 0.042; RR = 2.55*, P = 0.037* |
|                           | Harlozińska et al.,   | 314       | G1 versus G2/G3       | P = 0.0001                       | P = NS                           |
| Residual tumour size      | Chen et al., 1999     | 56        | ≥2 versus <2          | NA                              | RR = 0.96, P = 0.93              |
|                           | Harlozińska et al.,   | 314       | ≥2 versus <2          | P = 0.0001                       | P = NS                           |
| Lymph node involvement    | Tempfer et al., 1998  | 60        | Yes versus No         | RR = 2.8, P = 0.009             | RR = 2.8, P = 0.009              |
| Histological type         | Chen et al., 1999     | 56        | serous/mucinous versus others | NA                              | RR = 0.97, P = 0.73; RR = 1.04*, P = 0.7* |
|                           | Harlozińska et al.,   | 314       | serous versus others  | P = NS                           | P = NS                           |
| Age (years)               | Harlozińska et al.,   | 314       | ≥62 versus <62        | P = NS                           | P = NS                           |

*: Subset of 40 patients with residual tumour size ≤2 cm; NA: not available data; NS: non-significant statistical analysis.

From VEGF discovery till 2011, nine studies that directly correlated preoperative sVEGF with ovarian cancer outcome were published. Structured data extraction was performed on the articles to compare study populations, sVEGF assays, associations between sVEGF and clinicopathological characteristics, patient management, and outcome evaluation. Unfortunately, because of the heterogeneity of the studies and missing or incomplete information, it is not possible to pool data and to perform a meta-analysis in order to obtain univocal indications about sVEGF’s prognostic value.

The data reported in Tables 1, 3, and 4 show evident differences among studies. All studies included only epithelial ovarian carcinoma patients except those by Cooper et al. [25] and Li et al. [26], where other ovarian, peritoneal, and tubal malignancies were included. Patients underwent different chemotherapy regimens based on platinum analogues alone [21–24, 28] or in combination with taxane [25, 29]. Patients with early-stage disease were treated differently or were not treated, depending on the standards of the respective institutions. None of the studies exhaustively described followup (time, lost patients, events). Although seven [21–25, 27, 28] out of nine studies used the same sVEGF assay, one [22] of these measured significantly lower sVEGF values in ovarian cancer patients. Finally, widely differing sVEGF cut-off values (ranging from 100 to 826 pg/mL) were chosen for univariate and multivariate analysis, depending on the statistical methods chosen for the analysis.

In order to find out how sVEGF influences ovarian cancer biology, all studies analysed the association between sVEGF and the clinicopathological characteristics of the patients. The results seem to confirm that VEGF plays an important biological role in the pathogenesis of ascites [38, 39]. VEGF increases vessel permeability for circulating macromolecules,
thus facilitating extravasation of a plasma-rich exudate into the peritoneal cavity. Moreover, seven out of eight studies, concerning the relationship between sVEGF and FIGO stage, showed that VEGF concentrations measured in sera were not associated with FIGO stage. This may indicate that the effects promoted by VEGF are a continuous process and are independent of the clinical progression of the disease.

Interestingly, in our review of literature, sVEGF appears to be the best prognostic marker for OS in comparison with the established prognostic variables, since it stands out as an independent prognostic factor in most of the studies considered.

The scarcity of the data on the relationship between sVEGF levels and DFS makes it difficult to draw any firm conclusions in this regard. However, it is worth noting that sVEGF appears to be an independent prognostic factor for DFS in 2 out of 3 studies, as well as tumor stage and grade.

Chen et al. [23] and Hefler et al. [28] analyzed the prognostic value of sVEGF in a selected “low-risk” group of patients. Chen et al. [23] showed that sVEGF, FIGO stage and tumour grade were independent prognostic factors for OS and DFS in 40 patients with size tumour less than 2 cm. Hefler et al. [28] in a cohort of 56 patients with FIGO stage I found that sVEGF and tumour grade were independent prognostic factor for OS. The value of these results is conspicuous in those situations where the usefulness of adjuvant chemotherapy or the advisability of more chemotherapy cycles for certain categories of patients is under discussion.

In conclusion, almost all of the studies analysed in the present review, including the largest one by Hefler et al. [28], showed that elevated levels of sVEGF were significantly associated with shorter OS. It is worth noting that multiple phase III studies, ICON 7, GOG218, and OCEANS, have recently showed that the use of bevacizumab, a humanized antibody against VEGF, provides a clinically meaningful benefit in EOC patients outcome [40, 41].

Thus, from analysis of the literature reported in this review, as well as from results reported by recent clinical trials, sVEGF appears to be a promising prognostic factor in ovarian cancer that could identify a subgroup of patients with poor survival and higher risk of death that could benefit of bevacizumab therapy to improve their outcome.

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