Early outcome of a 31-gene expression profile test in 86 AJCC stage IB-II melanoma patients. A prospective multicentre cohort study

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Abstract
Background  The clinical and pathological features of primary melanoma are not sufficiently sensitive to accurately predict which patients are at a greater risk of relapse. Recently, a 31-gene expression profile (DecisionDx-Melanoma) test has shown promising results.

Objectives  To evaluate the early prognostic performance of a genetic signature in a multicentre prospectively evaluated cohort.

Methods  Inclusion of patients with AJCC stages IB and II conducted between April 2015 and December 2016. All patients were followed up prospectively to assess their risk of relapse. Prognostic performance of this test was evaluated individually and later combined with the AJCC staging system. Prognostic accuracy of disease-free survival was determined using Kaplan-Meier curves and Cox regression analysis. Results of the gene expression profile test were designated as Class 1 (low risk) and Class 2 (high risk).

Results  Median follow-up time was 26 months (IQR 22-30). The gene expression profile test was performed with 86 patients; seven had developed metastasis (8.1%) and all of them were in the Class 2 group, representing 21.2% of this group. Gene expression profile was an independent prognostic factor for relapse as indicated by multivariate Cox regression analysis, adjusted for AJCC stages and age.

Conclusions  This prospective multicentre cohort study, performed in a Spanish Caucasian cohort, shows that this 31-gene expression profile test could correctly identify patients at early AJCC stages who are at greater risk of relapse. We believe that gene expression profile in combination with the AJCC staging system could well improve the detection of patients who need intensive surveillance and optimize follow-up strategies. 

Received: 4 August 2018; Accepted: 4 January 2019

Conflicts of interest
The authors declare that they have no conflict of interest.

Funding source
Funding for this project was provided by Castle Biosciences, Inc. The research at the Melanoma Unit in Barcelona is partially funded by Spanish Fondo de Investigaciones Sanitarias grants 09/1393, 12/00840 and 15/00716; CIBER de Enfermedades Raras of the Instituto de Salud Carlos III, Spain, co-financed by European Development Regional Fund “A way to achieve Europe” ERDF; European Commission under the 6th Framework Programme, Contract No. LSHC-CT-2006-018702 (GenoMEL) and by the European Commission under the 7th Framework Programme, Diagnotics; The National Cancer Institute (NCI) of the US National Institute of Health (NIH) (CA83115), a grant from “Fundació La Marató de TV3, 201331-30”, Catalonia, Spain; CERCA Programme / Generalitat de Catalunya, and a grant from “Fundación Científica de la Asociación Española Contra el Cáncer”, Spain. Part of the work was carried out at the Esther Koplowitz Center, Barcelona.
Introduction
The incidence of melanoma is rapidly increasing and resulting mortality has risen significantly over the past 30 years.1 An increased incidence of invasive melanoma has been reported worldwide together with a corresponding rise in intermediate incidence of invasive melanoma has been reported.2–4 After the diagnosis of a primary melanoma, patients are enrolled in follow-up schedules depending on their risk of relapse, but due to its increasing prevalence, the workload and costs of surveillance programs have increased markedly.5–8 Many surveillance strategies have been proposed, mainly based on initial AJCC clinical–pathological staging;9–11 nevertheless, there are still many low-risk melanoma patients who suffer relapses, resulting in high mortality.9 In addition, approximately two thirds of patients whose melanoma will metastasize leading to death were initially sentinel lymph node biopsy (SLNB) negative (American Joint Committee on Cancer [AJCC] stages I and II).12,13
Due to this problem, many biomarkers have been developed to improve the identification of patients who will relapse, but with no real clinical impact until now.14 Recently, new prognostic tests based on genetic profiling signatures have been developed to better identify those patients who are at a higher risk of developing metastasis leading to death.15–20 A gene expression profile (GEP) test (DecisionDx-Melanoma, Castle Biosciences, Inc., Friendswood, TX, USA) evaluates 31 genes of primary cutaneous melanoma tissue and is independent from the clinical and pathological features of the tumour.15–24 However, this test has never before been evaluated in a prospective multicentre cohort.
Here we present a multicentre prospectively evaluated cohort of 86 patients where a 31-gene expression profile (GEP) test was used to assess their risk of relapse.

Material and methods
Study design
We conducted a prospective multicentre cohort study with patients from five tertiary melanoma referral centres in Spain included between April 2015 and December 2016, who were followed up until December 2018. Patients with resected pathologic American Committee on Cancer (AJCC, 7th edition, 2009) stages IB and II primary cutaneous melanoma were invited to participate in the study (Fig. 1). Furthermore, exclusion criteria included no evidence of disease within 3 months of primary surgery.
Patients were treated with conventional surgery followed by wide excision depending on Breslow thickness of the tumour. Patients with tumours T1b and above were staged with sentinel lymph node biopsy (SLNB) following each institution protocol. For the purpose of this study, patients with positive SLNB were excluded. The study was approved by the ethics committees of all Hospitals.

In all patients, tumour specimens were obtained from the hospital biobank. The formalin-fixed paraffin-embedded primary cutaneous melanoma tissue was analysed with the 31-GEP test (DecisionDx-Melanoma, Castle Biosciences, Inc).15,25 Moreover, the RT-PCR-based test classifies patients into a low-risk (Class 1) or high-risk (Class 2) category for recurrence, as previously reported and validated.15

Disease-free survival (DFS) was defined as the period of time in months from the date of diagnosis to the date of relapse. Patients that were free of disease at the time of the last follow-up or died during the study period, by any other causes, were treated as censored cases for evaluation purposes. All patients were followed up prospectively according to each institution protocol to identify relapses.

According to marked differences in 5-year survival curves for AJCC stages IIA and IIB patients (AJCC 79% vs. 68% respectively), all included subjects were further classified into two risk groups. Patients with AJCC stages IB and IIA were considered as low-risk and IIB and IIC as high-risk.18,26

For better reporting of the evaluation conducted, the REMARK checklist recommendations have been applied.

Statistical analysis
Pearson’s chi-squared and Student’s t-tests were used to compare categorical and continuous variables, respectively. Mann–Whitney–Wilcoxon was used to compare samples not distributed normally. Primary survival end-points (disease-free survival) were evaluated using Kaplan–Meier curves and univariate and multivariate Cox regression analyses. Due to the substantial censoring of survival times and several highly predictive covariates, we applied Firth’s Correction to our Cox regression model.27 The starting point for all cases was the diagnosis of
primary melanoma. P-values <0.05 were considered significant. Statistical analyses were performed using SPSS v.25.0 (IBM Corp., Armonk, NY, USA), and Cox regression analysis was performed using R studio (R Studio Team (2015). R Studio: Integrated Development for R. RStudio, Inc., Boston, MA, USA).

**Results**

After applying the inclusion/exclusion criteria, 88 patients were included in the study, but two were later excluded because of a technical failure in the test. Of the remaining 86 patients, 40 patients (46.5%) were male and 46 female (53.5%), with a median age at diagnosis of 59.2 years (interquartile range [IQR] 47–72). Characteristics of this cohort are summarized in Table 1. Patients were followed up for 2206 person-months, with a metastasis incidence rate of 3.17 cases per 1000 person-months. Overall median follow-up time was 26 months (IQR 22–30).

Class 1 and Class 2 patients showed a mean Breslow of 1.7 mm (standard deviation [SD] 1.4) and 3.7 mm (SD 2.9), respectively, which was statistically significant (P < 0.001). Moreover, Class 2 melanomas were ulcerated more often and presented significantly higher AJCC staging (Table 1).

Relapses were identified in seven patients (8.1%), all corresponding to Class 2 (high-risk) by the GEP test (P < 0.001). Furthermore, the GEP risk score identified 19 patients (22.1%) with a risk score different from that predicted by AJCC classification. Five (5.8%) patients with high-risk AJCC stage were rated as Class 1 (low risk), and 14 subjects (16.3%) with low-risk AJCC stage were identified as Class 2 (high risk) by the GEP test. Five patients (5.8%) presented relapses with a high-risk GEP test score and AJCC high-risk at the same time, while two subjects (2.3%) patient were identified as Class 2 (high-risk) by the GEP test although belonging to AJCC low-risk.

Kaplan–Meier survival curves were analysed to evaluate DFS for the two 31-GEP risk classes and showed statistically significant differences between the two groups (log rank P < 0.001). When combining the GEP test with the AJCC (log rank P = 0.001) the significance was maintained (Fig. 2).

Following curve comparison by Kaplan–Meier method, we compared the prognostic accuracy of the GEP test to the AJCC and age using Cox regression analysis. Both univariate and multivariate analysis showed the GEP test to be an independent predictor of metastasis. Hazard ratios for the GEP test were 28.37 (95% CI 3.46–3682.91; P < 0.01) for the univariable analysis and 18.82 (95% CI 1.81–2549.76; P = 0.01) for the multivariable analysis (Table 2).

**Discussion**

This paper describes the results of a 31-GEP test performed to better categorize patients with low to intermediate-risk melanoma according to the AJCC staging system. We present the data from a multicentre cohort study in which patients with malignant melanoma were followed up prospectively after the

| Table 1 Basal clinicopathological characteristics of the cohort and relapse data |
|-----------------------------|----------------|----------------|----------------|-------------|
|                             | Overall        | Class 1 Low risk | Class 2 High risk | P value    |
|------------------------------|----------------|----------------|-----------------|-------------|
| n                            | 86             | 53             | 33              |             |
| Follow-up time median (IQR)  | 26 (22−30)     | 27 (23−32)     | 24 (20−29)      | 0.066       |
| Sex                          |                |                |                 |             |
| Male                         | 40 (46.5%)     | 21 (40%)       | 19 (58%)        | 0.105       |
| Female                       | 46 (53.5%)     | 32 (60%)       | 14 (42%)        |             |
| Age                          |                |                |                 |             |
| Median (IQR)                 | 59.2 (47–72)   | 57 (46–68)     | 68 (55–74)      | 0.025       |
| Localization                 |                |                |                 |             |
| Acral                        | 5 (6%)         | 2 (4%)         | 3 (9%)          | 0.738       |
| Head and neck                | 11 (13%)       | 7 (13%)        | 4 (12%)         |             |
| Legs                         | 21 (24%)       | 15 (28%)       | 6 (18%)         |             |
| Arms                         | 12 (14%)       | 7 (13%)        | 5 (15%)         |             |
| Trunk                        | 37 (43%)       | 22 (41%)       | 15 (45%)        |             |
| Breslow                      |                |                |                 |             |
| Mean (SD)                    | 2.5 (2.3)      | 1.7 (1.4)      | 3.7 (2.9)       | <0.001      |
| <1.00 mm                     | 18 (21%)       | 17 (32%)       | 1 (3%)          | <0.001      |
| 1.00–2.00 mm                 | 33 (38%)       | 23 (43%)       | 10 (30%)        |             |
| 2.01–4.00 mm                 | 22 (26%)       | 10 (19%)       | 12 (36%)        |             |
| >4.00 mm                     | 13 (15%)       | 3 (6%)         | 10 (30%)        |             |
| Mitotic rate (mm²)           |                |                |                 |             |
| <1 mm²                       | 9 (11%)        | 6 (11%)        | 3 (9%)          | 0.684       |
| ≥1 mm²                       | 76 (88%)       | 46 (87%)       | 30 (91%)        |             |
| N.A.                         | 1 (1%)         | 1 (2%)         | 0               |             |
| Ulceration                   |                |                |                 |             |
| Absent                       | 60 (70%)       | 51 (96%)       | 9 (27%)         | <0.001      |
| Present                      | 26 (30%)       | 2 (4%)         | 24 (73%)        |             |
| AJCC stage                   |                |                |                 |             |
| Low-risk (IIB-IIA)           | 62 (72%)       | 48 (91%)       | 14 (42%)        | <0.001      |
| High-risk (IIB-IIIC)         | 24 (28%)       | 5 (9%)         | 19 (58%)        |             |
| Relapse                      |                |                |                 | <0.001      |
| No                           | 79 (91%)       | 53 (100%)      | 26 (79%)        |             |
| Yes                          | 7 (9%)         | 0              | 7 (21%)         |             |
| Relapse site (n − 7)         |                |                |                 |             |
| Skin                         | 0              | 2 (29%)        | –               |             |
| Lymph node                   | 0              | 2 (29%)        | –               |             |
| Visceral                     | 0              | 3 (43%)        | –               |             |

AJCC, American Joint Committee on Cancer; IQR, Interquartile range; N.A., Not Available; SD, Standard Deviation. P-values are bold where they are less than or equal to the significance level cut-off of 0.05.

31-GEP test. We observed that seven patients (8.1%) presented relapses within a median of 12 months (IQR 5–21), all seven belonging to the Class 2 group (overall, 21.2% of the Class 2 group). Kaplan–Meier survival curves were statistically different between both groups with an increased risk of recurrence in the Class 2 group with a hazard ratio of 28.37 and 18.82, for univariate and multivariate Cox regression analysis, respectively. Similarly, a previous validation study of the 31-GEP test showed that the metastatic risk was predicted with high accuracy in the Class
2 cohort of primary cutaneous melanomas, with a receiver operating characteristic (ROC) curve value of 0.91–0.93.15 Subsequent studies have analysed the use of the 31-GEP test in combination with the AJCC stage system and found that this approach improves the identification of patients at risk of relapse.18,19 In this report, we also combined GEP with the AJCC score confirming that patients with higher AJCC (IIB-IIC) staging and Class 2 GEP are at a greater risk of relapse. Nevertheless, two patients with a low-risk AJCC stage and a Class 2 GEP metastasized. Based solely on AJCC staging, many patients classified at low-risk will relapse as seen in the study by Ferris et al. where 43% of the cases classified as Class 2 GEP and AJCC low risk, relapsed. This has a profound impact on high-risk patients based on the GEP test, but at lower AJCC stages who could opt for intensive follow-up,9 but are at early stages of the AJCC. Moreover in the new era of adjuvant therapies, patients at greater risk of relapse could be selected for these treatment schemes and improve overall survival.

Moreover, GEP tests have been used in combination with SLNB status resulting in different outcomes of disease-free survival, and overall survival curves.16 The authors have observed that patients with negative SLNB and class 2 GEP have a worse prognosis than patients with positive SLNB and class 1 GEP status.16,21 In the present study, we only included SLN negative patients, to evaluate the performance of this test in identifying those at a high risk of melanoma relapsing during early stages. It may be that the use of genetic signatures in conjunction with the classic clinical, pathological criteria of AJCC and the SLNB status will significantly improve the detection of patients at risk of relapse. Accordingly, we believe that validating a gene assay test will be an important step towards the goal of optimizing follow-up protocols and we propose personalized adjuvant therapy in low to intermediate-risk melanoma patients.

As a limitation of this study, we observed that patients who tested as Class 2 GEP had a tendency towards thicker
melanomas, higher frequency of ulceration and higher AJCC stages. However, univariate and multivariate Cox regression analyses showed that the GEP test was an independent prognostic factor. Moreover, we had a low incidence of events, so the external validity should be interpreted with caution. This could be explained by the follow-up time being relatively short, although most of the metastases developed within the first two years of follow-up. The number of relapses is then, in accordance with that published in the literature.\(^6,13\) Moreover, due to the multicentre and prospective design of this study and the assessment of this test in 'real daily practice', the GEP test gives useful information about what to expect during the first two years of follow-up, as we observed that 21% of the patients in the class 2 group developed metastasis.

**Conclusion**

We observed that a 31-GEP test allows the accurate prediction of patients at a high risk of relapse, despite having a low-risk AJCC staging. We believe that GEP in combination with AJCC staging system could well improve the detection of patients who need intensive surveillance and optimize follow-up strategies.

**Acknowledgements**

Thanks to our patients and their families who are the main reason for our studies; to nurses from the Melanoma Unit of Hospital Clinic of Barcelona, Daniel Gabriel, Pablo Iglesias and Maria E Moliner for helping to collect patient data and to Paul Hetherington for helping with English editing and correction of the manuscript.

**Role of the sponsors**

The sponsors had no role in the design and conduct of the study, nor in the collection, analysis and interpretation of data, nor in the preparation, review and approval of the manuscript nor in the decision to submit the manuscript for publication.

**Ethical approval**

All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards.

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