The surgical treatment of non-metastatic melanoma in a Clinical National Melanoma Registry Study Group (CNMR): a retrospective cohort quality improvement study to reduce the morbidity rates

Antonella Vecchiato, Simone Mocellin, Paolo Del Fiore, Giulio Tosti, Paolo A. Ascierto, Maria Teresa Corradin, Vincenzo De Giorgi, Giuseppe Giudice, Paola Queirolo, Caterina Ferrelli, Marcella Occelli, Monica Giordano, Giusto Trevisan, Luigi Mascheroni, Alessandro Testori, Romina Spina, Alessandra Buja, Francesco Cavallini, Corrado Caraco, Antonio Sommariva, Carlo Riccardo Rossi on behalf of the Clinical National Melanoma Registry Study Group at the Italian Melanoma Intergroup (IMI)

Abstract

Background: Reproducible, high-quality surgery is a key point in the management of cancer patients. Quality indicators for surgical treatment of melanoma has been presented with benchmarks but data on morbidity are still limited. This study presents the quality indicators on morbidity after surgical treatment for non-metastatic skin melanoma in an Italian registry.

Methods: Data were extracted from the Central National Melanoma Registry (CNMR) promoted by the Italian Melanoma Intergroup (IMI). All surgical procedures (WE, SLNB or LFND) for non-metastatic skin melanoma between January 2011 and February 2017 were evaluated for inclusion in the study. Only centers with adequate completeness of information (> 80%) were included in the study. Short-term complications (wound infection, dehiscence, skin graft failure and seroma) were investigated.

Results: Wound infection rate was 1.1% (0.4 to 2.7%) in WE, 1.3% (0.7 to 2.5%) in SLNB and 4.1% (2.1 to 8.0%) in LFND. Wound dehiscence rate was 2.0% (0.8 to 5.1%) in WE, 0.9% (0.2 to 3.0%) in SLNB and 2.8% (0.9 to 8.6%) in LFND. Seroma rate was 4.2% (1.5 to 11.1%) in SLNB and 15.1% (4.6 to 39.9%) in LFND. Unreliable information was found on skin graft failure.

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Background

Worldwide, 287,723 new cases of cutaneous melanoma are diagnosed each year, causing the death of 60,712 patients [1]. In non-metastatic melanoma (i.e. without clinically evident regional lymph node or distant metastasis), surgery is the mainstay of treatment [2]. The surgical strategy involves a combination of wide excision (WE), sentinel lymph node biopsy (SLNB) and radical lymph node dissection (LFND), according to cancer staging. WE can be virtually part of the surgical management of all patients with skin melanoma, SNLB is recommended based on the primary tumor thickness and LFND may be required in patients with positive sentinel node [3].

Reproducible, high-quality surgery is a key point in the management and prognosis of cancer patients [4, 5]. When a standardized surgical procedure is established, a cyclic audit (including collection, analysis and feedback of both procedural and outcome data) allows for monitoring and improving the quality of surgery. However, cancer surgery is not always standardized, thus limiting the application of such approach [6].

While quality of surgery has been already investigated in other cancers, the use of measures of quality assurance for surgery is less established in melanoma [6]. Available literature offers substantial heterogeneity in surgical procedures among melanoma centers or even among surgeons within the same center [7]. Although adherence to current standards is part of a quality assurance process, the spreading of clinical practice guidelines is not sufficient per se to warrant homogeneity and quality of surgical treatment [8]. There is a growing interest in the implementation of a quality assurance program that includes a quantitative analysis of a set of quality indicators, such as those that can be extracted from electronic medical records [6].

Since 2014, the Italian Melanoma Intergroup (IMI) has been promoting the standardization and quality control of surgical treatment of stage I-III melanoma in Italy [9]. The IMI achieved an expert consensus on surgical treatment of melanoma, defining a list of quality indicators (detection rate, false negative rate, minimum number of excised lymph nodes, postoperative morbidity and local recurrence) with reference to benchmark values, which could be the basis for a standardized quality assurance program in Italy [10]. While quality indicators for detection rate, false negative rate and minimum number of excised lymph nodes were presented for the Central National Melanoma Registry (CNMR), data on morbidity were still pending and data on recurrence are waiting for adequate follow-up [10].

The present study focuses on evaluation of the quality indicators about morbidity after surgical treatment for non-metastatic skin melanoma.

Methods

This is a prospective multicenter study on postoperative morbidity after surgical treatment for non-metastatic skin melanoma. Data were extracted from the Central National Melanoma Registry (CNMR), which is a prospectively maintained national database for melanoma treatment promoted by the IMI in 43 melanoma centers in Italy [9]. The study was conducted according to Helsinki Declaration principles and was approved by the Ethics Committee of the Central National Melanoma Registry (CNMR). All patients gave their consent to have their data collected for scientific purpose.

The following short-term (within 30 days after surgery) complications were investigated: infection, dehiscence, skin necrosis after WE; infection, dehiscence and seroma after SLNB or LFND. Complications were defined using the National Cancer Institute Common Terminology Criteria for Adverse Events v4.0 [11]. Wound infection was defined as a disorder characterized by an infectious process involving the wound. Wound dehiscence was defined as a finding of separation of the approximated margins of a surgical wound. Skin necrosis was defined as a death of cells in living tissue caused by external factors such as infection, trauma, or bacterial agents. Seroma was defined as finding of a collection of serum in the tissues. Patient demographics were also collected.

All surgical procedures (WE, SNLB or LFND) for non-metastatic skin melanoma between January 2011 and February 2017 were evaluated for inclusion in the study. Only centers with adequate completeness of information (> 80%) were included in the study (11 centers for analysis of morbidity after WE; 14 centers for analysis of morbidity after SNLB; 11 centers for analysis of morbidity after LFND). Within selected centers, patients with...
pending information on morbidity and those with un-
filled form were excluded from the analysis. Patient se-
lection is shown in Supplementary Figs. 1, 2 and 3.

Statistical analysis was conducted using R 3.5 (R Foun-
dation for Statistical Computing, Vienna, Austria) [12].
Categorical data were expressed as number and per-
centage, and continuous data as median and interquar-
tile range (IQR). The morbidity rate was pooled among cen-
ters using a generalized linear mixed-effects model (with
the center as random effect). The I² value was used to
investigate heterogeneity, and meta-regression was used
to explore the role of center size and groin site on het-
erogeneity. A p-value less than 0.05 was considered sta-

tistically significant.

Results

Wide excision
The analysis included 3118 patients (1647 males and
1471 females) who underwent WE in 11 centers (median
246 patients/center; min 90 patients/center, max 650 pa-

tients/center). Median age was 57 years (IQR 45–70) and
median BMI was 24.8 kg/m² (IQR 22.4–27.5). Comor-

bidity was present in 459 patients (15.1%), while the
information was not available in 80 patients. Overall,
median Charlson Comorbidity Index was 2 (IQR 0–3).

Wound infection occurred in 44 patients and dehis-
cence in 100. The information on skin graft failure was
not reliable due to the non-negligible number of unfilled
forms (21%).

The proportion of patients developing wound infection
ranged from 0.002 to 0.167 among centers, with a
pooled proportion of 0.011 (95% CI 0.004 to 0.027)
(Fig. 1). Heterogeneity was high (I² = 87%) and center
size (> 200 vs. < 200 patients) was found to have a sig-
nificant contribution to heterogeneity (p < 0.0001).
Summary estimates of proportion of patients developing
wound infection were 0.051 (95% CI 0.021 to 0.121; I² = 80%)
in centers with low size (< 200 patients) and 0.006
(95% CI 0.003 to 0.009; I² = 0%) in centers with large
size (> 200 patients).

The proportion of patients developing dehiscence ran-
ded from 0.005 to 0.533 among centers, with a pooled
proportion of 0.020 (95% CI 0.008 to 0.051) (Fig. 1). Heter-
ogeneity was high (I² = 93%) and center size (> 100 vs. <
100 patients) was found to have a significant contribu-
tion to heterogeneity (p < 0.0001). Summary estimate of pro-
portion of patients developing dehiscence was 0.016 (95% CI
0.010 to 0.024; I² = 54%) in centers with large size (> 100
patients) and 0.533 (95% CI 0.425 to 0.639) in the only
center with low size (< 100 patients).

Sentinel lymph node biopsy
The analysis included 1853 patients (993 males and 860
females) who underwent SLNB in 14 centers (median 98
patients/center; min 24 patients/center, max 338 pa-
tients/center). SLNB site was axilla (1058 patients,
57.1%), neck (163 patients, 8.8%) or groin (632 patients,
34.1%). Median age was 57 years (IQR 45–69) and me-
dian BMI was 25.1 kg/m² (IQR 22.7–28.0). Comorbid-
ities were present in 304 patients (16.7%), while the
information was not available in 40 patients. Overall,
median Charlson Comorbidity Index was 1 (IQR 0–3).

Early wound complications occurred in 176 patients,
including 29 wound infections, 39 dehiscences and 139
seromas (not mutually exclusive).

The proportion of patients developing early wound
complications ranged from 0.007 to 0.831 among cen-
ters. Summary estimate of proportion of patients devel-
oping early wound complications after SLNB was 0.069
(95% CI 0.030 to 0.150) (Fig. 2). Heterogeneity was high
(I² = 94%); neither center size (p = 0.11) or rate of groin
sites among SLNBs (p = 0.84) were not found to have a
significant contribution to heterogeneity.

The proportion of patients developing wound infection
ranged from 0.005 to 0.104 among centers. Summary
estimate of proportion of patients developing wound
infection after SLNB was 0.013 (95% CI 0.007 to 0.025)
(Fig. 2). Heterogeneity was moderate (I² = 60%); neither
center size (p = 0.25) or rate of groin sites among SLNBs
(p = 0.60) were not found to have a significant contribu-
tion to heterogeneity.

The proportion of patients developing dehiscence ran-
ded from 0.005 to 0.101 among centers. Summary es-
estimate of proportion of patients developing dehiscence
after SLNB was 0.009 (95% CI 0.002 to 0.030) (Fig. 2).
Heterogeneity was high (I² = 88%); neither center size
(p = 0.98) or rate of groin sites among SLNBs (p = 0.89)
were not found to have a significant contribution to hetero-

Radical lymph node dissection
The analysis included 502 patients (301 males and 201
females) who underwent LFND in 11 centers (median
25 patients/center; min 15 patients/center, max 138 pa-
tients/center). LFND site was axilla (276 patients,
55.0%), neck (60 patients, 12.0%) or groin (166 patients,
33.0%). Median age was 59 years (IQR 47–70) and me-
dian BMI was 25.5 kg/m² (IQR 22.9–28.6). Comorbid-
ities were present in 93 patients (19.0%), while the
information was not available in 12 patients. Overall,
median Charlson Comorbidity Index was 2 (IQR 0–3).
Early wound complications occurred in 98 patients, including 22 wound infections, 27 dehiscences and 85 seromas (not mutually exclusive).

The proportion of patients developing early wound complications ranged from 0.024 to 0.969 among centers. Summary estimate of proportion of patients developing early wound complications after LFND was 0.195 (95% CI 0.068 to 0.447) (Fig. 3). Heterogeneity was high ($I^2 = 94%$); neither center size ($p = 0.08$) or rate of groin sites among LFNDs ($p = 0.41$) were not found to have a significant contribution to heterogeneity.

The proportion of patients developing wound infection ranged from 0.006 to 0.58 among centers. Summary estimate of proportion of patients developing wound infection after SLNB was 0.041 (95% CI 0.021 to 0.080) (Fig. 3). Heterogeneity was moderate ($I^2 = 54%$); neither center size ($p = 0.07$) or rate of groin sites among SLNDs ($p = 0.53$) were not found to have a significant contribution to heterogeneity.

The proportion of patients developing dehiscence ranged from 0.006 to 0.244 among centers. Summary estimate of proportion of patients developing dehiscence after SLNB was 0.028 (95% CI 0.009 to 0.086) (Fig. 3). Heterogeneity was high ($I^2 = 80%$); neither center size ($p = 0.38$) or rate of groin sites among SLNDs ($p = 0.14$) were not found to have a significant contribution to heterogeneity.

The proportion of patients developing seroma ranged from 0.021 to 0.969 among centers. Summary estimate of proportion of patients developing seroma after SLNB was 0.151 (95% CI 0.046 to 0.399) (Fig. 3). Heterogeneity was high ($I^2 = 95%$); neither center size ($p = 0.18$) or rate of groin sites among SLNDs ($p = 0.29$) were not found to have a significant contribution to heterogeneity.

**Discussion**

This study reported the morbidity rates in the largest series of non-metastatic melanoma patients in Italy and one of the largest series worldwide. The present work allowed to evaluate standardization and quality of surgical treatment of cutaneous melanoma within the frame of the Italian Melanoma Intergroup, which is the largest Italian scientific organization dedicated to the management of patients with this disease [10].

Interestingly, we found lower occurrence of wound infection after WE or LFND when compared to available literature [13] (Table 1), thus suggesting limited opportunities for further improvements now. Such rates can be used as morbidity benchmarks in addition to available quality indicators for surgical treatment of melanoma [10].

Interestingly, we found lower occurrence of wound infection after WE or LFND when compared to available literature [13] (Table 1). On one hand, this difference might be partially due to different definitions of wound infection, i.e. presence of fever, only skin redness, suppuration versus cellulitis, isolation of bacteria from wound. On the other hand, some factors might have led
to such difference, including: i) recording of infections occurring only during hospital stay or also at the time of outpatient department visits; ii) different frequency/type of antibiotic prophylaxis; iii) different surgical techniques (i.e., use, type and duration of permanence of drainages in the surgical wound).

Postoperative infections account for around one out of four complications associated with hospital-related health care procedures, and can impair patient prognosis [16]. Beyond clinical importance, a low infection rate can also have an economic impact, since the management of hospital-related infections requires about 0.8% of gross domestic product (GDP) in Italy [17].

Unfortunately, the information on skin graft failure was not reliable because one out of five forms did not report such complication. This situation likely occurred because skin graft failure required a reconstructive
surgical management that several centers demanded to a different surgical unit. This situation may be addressed by improving the information exchange among surgical centers involved in patient care.

Of note, morbidity rates showed high heterogeneity across melanoma centers, underlying the role of the center itself on this matter. Our data suggested an association between higher morbidity rate and small-volume centers, thus confirming the relationship between patient outcome and hospital surgical volume [18].

The present study contributes to the definition of quality indicators for surgical treatment for non-metastatic skin melanoma, by adding morbidity indicators that can be used as the basis for a standardized quality assurance program [10]. The importance of this topic relies on the large number of surgical

| Center size | Proportion [95% CI] |
|-------------|---------------------|
| 15          | 0.969 [0.782, 1.000] |
| 18          | 0.111 [0.014, 0.347] |
| 19          | 0.474 [0.244, 0.711] |
| 21          | 0.048 [0.001, 0.238] |
| 23          | 0.043 [0.001, 0.219] |
| 25          | 0.200 [0.068, 0.407] |
| 45          | 0.333 [0.200, 0.490] |
| 45          | 0.467 [0.317, 0.621] |
| 68          | 0.338 [0.228, 0.463] |
| 85          | 0.024 [0.003, 0.082] |
| 138         | 0.029 [0.008, 0.073] |

**RE Model**

- Early wound complications: 0.195 [0.068, 0.447]
- Wound infection: 0.041 [0.021, 0.080]
- Dehiscence: 0.028 [0.009, 0.086]
- Seroma: 0.151 [0.046, 0.399]

Fig. 3 Patients developing early wound complications, wound infection, dehiscence and seroma after LFND: forest plot
procedures for non-metastatic skin melanoma [2], thus patient management and prognosis can benefit from quality control and standardization of such procedures [4, 5].

The strengths of this study included the prospective collection of data in a national registry, the multicenter design and the standardized definitions of the complications [11]. The study has some limitations. First, a considerable number of centers were excluded due to poor completeness of data. Although this approach allowed limiting the impact of low-quality data on the study results, the representativeness of the included centers may be limited. Future developments will aim to achieve adequate completeness of data in the excluded centers and will implement regular audits. Second, the occurrence on skin graft failure after WE could not be evaluated due to the non-negligible number of unfilled forms. This limitation can be addressed by future improvements regarding information exchange among surgical centers involved in patient care.

Conclusion
Our findings contribute to available literature in setting up the recommended standards for melanoma centers, thus improving the quality of surgery offered to patients. Such quality indicators can be used by other hospitals to direct quality improvement efforts.

Table 1 Referral values for morbidity rate in the IMI-CNMR study and in the international

| Surgical Procedure | Indicators                  | Benchmark referral values |
|--------------------|-----------------------------|---------------------------|
|                    | Present study               | International literature  |
| WE                 | Wound infection             | 1.1% (0.4 to 2.7%)        | 4.6–8.4%a                  |
|                    | Wound dehiscence            | 2.0% (0.8 to 5.1%)        | 3.5–4.6%a                  |
|                    | Skin graft failure          | unreliable                | < 2%a                     |
| SLNB               | Wound Infection             | 1.3% (0.7 to 2.5%)        | 2.9% (1.5 to 4.6%)b        |
|                    | Wound dehiscence            | 0.9% (0.2 to 3.0%)        | 0.24–1.2%a                 |
|                    | Seroma                      | 4.2% (1.5 to 11.1%)       | 5.1% (2.5 to 8.6%)b        |
| LFND               | Wound infection             | 4.1% (2.1 to 8.0%)        | 15.8%a                     |
|                    | Wound dehiscence            | 2.8% (0.9 to 8.6%)        | 3%a                       |
|                    | Wound infection and/or dehiscence | 6.5% (2.9 to 14.0%)    | 21.6% (13.8 to 30.6%)c     |
|                    | Seroma                      | 15.1% (4.6 to 39.9%)      | 17.9% (10.3 to 27%)c        |

Data expressed as percentage with 95% confidence interval in parentheses

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Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s12885-020-07705-4.

Additional file 1: Supplementary Figure 1. Flow-chart of patient inclusion in wide excision (WE) analysis. Supplementary Figure 2. Flow-chart of patient inclusion in sentinel lymph node biopsy (SLNB) analysis. Supplementary Figure 3. Flow-chart of patient inclusion in radical lymph node dissection (LFND) analysis.

Abbreviations
WE: Wide excision; SLNB: Sentinel lymph node biopsy; LFND: Radical lymph node dissection; IMI: Italian Melanoma Intergroup; CNMR: Clinical National Melanoma Registry; GDP: Gross domestic product

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^Clinical National Melanoma Registry Study Group at the Italian Melanoma Intergroup:
Asero Salvatore: Azienda Ospedaliera di Rilievo Nazionale e di Altra Specializzazione Garibaldi-Nesima, Catania, Italy.
Barbati Rosanna: ASL Roma C - Ospedale S. Eugenio, Roma, Italy.
Bianchi Luca: Azienda Ospedaliera Universitaria Policlinico Tor Vergata, Roma, Italy.
Bruder Francesca: Ospedale Oncologico Cagliari, Cagliari, Italy.
Catricala Caterina: Istituto Dermatologico San Gallicano IRCCS – IFO, Roma, Italy.
Cineri Saverio: Presidio Ospedaliero Antonio Perrino, Brindisi, Italy.
Del Vecchio Michele: Fondazione I.R.C.C.S. Istituto Nazionale dei Tumori, Milano, Italy.
Di Filippo Franco: Istituto Nazionale Tumori Regina Elena IRCCS – IFO, Roma, Italy.
Fargnoli Maria Concetta: Presidio Ospedaliero San Salvatore, L’Aquila, Italy.
Fierro Maria Teresa: A.O.U. Città della Salute e della Scienza - P.O. San Lazzaro, Torino, Italy.
Forcignano Rosachira: Azienda Ospedaliera Vito Fazzi, Lecce, Italy.

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| SLNB               | Wound Infection             | 1.3% (0.7 to 2.5%)        | 2.9% (1.5 to 4.6%)b        |
|                    | Wound dehiscence            | 0.9% (0.2 to 3.0%)        | 0.24–1.2%a                 |
|                    | Seroma                      | 4.2% (1.5 to 11.1%)       | 5.1% (2.5 to 8.6%)b        |
| LFND               | Wound infection             | 4.1% (2.1 to 8.0%)        | 15.8%a                     |
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|                    | Seroma                      | 15.1% (4.6 to 39.9%)      | 17.9% (10.3 to 27%)c        |
GUIDOBONI MASSIMO: IR.S.T. Istituto Scientifico Romagnolo per lo Studio e la Cura dei Tumori, Meldola, Italy.
IACONO CARMELINO: Azienda Ospedaliera Sanità 7 Ragusa - Ospedale Maria Paternò, Arezzo, Italy.
LOSPALLUTI LUCIA: Azienda Sanitaria Locale BA - Ospedale di Venere, Bari, Italy.
MAIO MICHELE: Azienda Ospedaliera Universitaria Senese Ospedali Le Scoote, Siena, Italy.
MILES LAURA: Azienda Ospedaliera Papa Giovanni XXIII, Bergamo, Italy.
MOISE GIANMICHELE: Azienda per i Servizi Sanitari n°2 Isontina Ospedale di Gorizia Dipartimento di Medicina, Gorizia, Italy.
MORETTI GIOVANNA: Azienda Ospedaliera Ospedali Riuniti Papardo–Piemonte, Messina, Italy.
PELLICINO RICCARDO: IRCCS Casa Sollievo della Sofferenza, San Giovanni Rotondo, Italy.
PIZZICHELLA MARIA ANTONIETTA: Centro di Riferimento Oncologico, Istituto Nazionale Tumori, Aviano, Italy.
RINALDI GAETANA: Azienda Ospedaliera Universitaria Policlinico “Pao Giaccone, Palermo, Italy.
SANNI GIOVANNI: Azienda Ospedaliero-Universitaria di Sassari, Sassari, Italy.
STAIBANO STEFANIA: Azienda Ospedaliera Universitaria Federico II di Napoli, Napoli, Italy.
VISINI MARLENA: A.O. di Lecco Presidio Ospedaliero Alessandro Manzoni, Lecco, Italy.
ZANNETTI GUIDO: Azienda Ospedaliero-University of Bologna Policlinico S. Orsola -Malpighi, Bologna, Italy.
ZICCHICI LEONARDO: Azienda Sanitaria Provinciale - Presidio Ospedaliero di Triapani, Trapani, Italy.

Authors’ contributions
Study concepts: PDF, FC, SM, AV. Study design: PDF, FC, SM, AV, CRR. Data acquisition: PDF, FC, RS, MTC, GT, VDG, GG, PQ, MG, CF, GT, LM, MO, AT, IS, CC, PA. Quality control of data and algorithms: PDF, FC, SM, AS. Statistical analysis: FC. Manuscript preparation: PDF, FC, SM, AS. Manuscript editing: PDF, FC, SM. Manuscript review: SM, AB, CRR, MC, PA. Data analysis and acquisition: PDF, FC, RS, MTC, GT, VDG, GG, PQ, MG, CF, GT, LM, MO, AT, IS, CC, PA. Quality control of data and algorithms: PDF, FC, SM, AV. Study design: PDF, FC, SM, AV, CRR. Data acquisition: PDF, FC, RS, MTC, GT, VDG, GG, PQ, MG, CF, GT, LM, MO, AT, IS, CC, PA. Quality control of data and algorithms: PDF, FC, SM, AS. Statistical analysis: FC. Manuscript preparation: PDF, FC, SM, AS. Manuscript editing: PDF, FC, SM. Manuscript review: SM, AB, CRR. All authors contributed to manuscript revision, read, and approved the submitted version.

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Availability of data and materials
The datasets generated and analysed during the current study are available on Mendeley Data.

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Ethics approval and consent to participate
The study was approved by the Ethics Committee of Veneto Institute of Oncology (approval No. CESC-IOV 2011/17) on 03 February 2015. All procedures performed in studies involving human participants were in accordance with the 1964 Helsinki declaration and its later amendments or comparable ethical standards. Written informed consent was obtained from all individual participants included in the study.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

Author details
1Surgical Oncology Unit, Veneto Institute of Oncology IOV – IRCCS, Padova, Italy. 2Department of Surgery, Oncology and Gastroenterology (DISCOG), University of Padua, Padova, Italy. 3Division of Melanoma, Sarcoma and Rare Tumors, IRCCS, European Institute of Oncology, Milan, Italy. 4Department of Melanoma and Cancer Immunotherapy, Istituto Nazionale Tumori IRCCS Fondazione Pascale, Naples, Italy. 5Department of Dermatology, Santa Maria degli Angeli Hospital, Pordenone, Italy. 6Section of Dermatology, Department of Health Sciences, University of Florence, Florence, Italy. 7Division of Plastic and Reconstructive Surgery and Burn Unit, University of Bari, Bari, Italy. 8Division of Medical Oncology for Melanoma, Sarcoma, and Rare Tumors, IEO, European Institute of Oncology IRCCS, Milan, Italy. 9Section of Dermatology, Department of Medical Sciences and Public Health, University of Cagliari, Cagliari, Italy. 10Medical Oncology Unit, Santa Croce and Carle Teaching Hospital, Cuneo, Italy. 11Pathology, ASST-Lariana, Ospedale Sant’Anna, Como, Italy. 12DSM-Department of Medical Sciences, University of Trieste, Trieste, Italy. 13Unit of General Surgery, San Pio X Hospital, Milan, Italy. 14Department of Dermatology, Fondazione IRCCS Policlinico San Matteo, 27100 Pavia, Italy. 15Department of Cardiological, Thoracic and Vascular Sciences, and Public Health, University of Padova, Padova, Italy. 16Independent Statistician, Solagna, Italy. 17National Cancer Institute Fondazione G. Pascale, SC Chirurgia Melanoma e dei Tumori Cutanei, Naples, Italy. 18Unit of Surgical Oncology of the Esophagus and Digestive Tract, Veneto Institute of Oncology IOV-IRCCS, Padua, Italy.

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