This decade has brought significantly improved outcomes for patients with advanced melanoma with immunotherapies and targeted treatments offering utility in a variety of settings. In 2020, we can hope for durable long-term responses, and complete remission in a subset of patients with metastatic disease. In the adjuvant setting, approximately 50% improvements in recurrence-free survival are seen both with targeted and immunotherapies. Early data from neoadjuvant immunotherapy clinical trials are very promising. However, responses to treatment are heterogeneous and not always durable; further advances are required, and several emerging strategies are of particular interest. We review the systemic treatment of melanoma, discussing the treatment of unresectable stage III–IV and recurrent disease, outlining curative treatment of cutaneous melanoma in the adjuvant setting and briefly discussing neoadjuvant systemic therapies for advanced melanoma.

Key words: melanoma; systemic therapy; targeted therapy; immunotherapy.

Accepted Apr 27, 2020; Epub ahead of print Apr 28, 2020
Acta Derm Venereol 2020; 100: adv00141.
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Accounting for only 1% of all skin malignancies, melanoma represents the most aggressive and deadly form of skin cancer (1). Melanoma is predominantly a disease of Caucasian populations and affects men and women in equal measure. With a propensity to migrate to draining lymph nodes and any visceral organ, metastatic melanoma carries a poor prognosis.

Prior to 2011, outcomes were poor, with treatment for metastatic disease limited to palliative therapies that offered little or no survival benefit. In 2020, we can hope for durable long-term responses, and complete remission in a subset of patients. The use of immunotherapies and targeted therapies for melanoma in the metastatic, adjuvant and neoadjuvant settings will be reviewed here; the initial management of cutaneous melanoma is discussed separately. This review will cover the systemic treatment of melanoma, starting with a description of therapeutic agents. We will discuss the treatment of unresectable stage III–IV and recurrent disease, outlining curative treatment of cutaneous melanoma in the adjuvant setting and briefly discuss neoadjuvant systemic therapies for advanced melanoma.

SIGNIFICANCE
Melanoma is an aggressive and rare skin cancer that can threaten the lives of patients it affects. New treatments have been introduced over the past decade which have dramatically changed the way in which patients with advanced melanoma are managed. Here we review the treatments currently available to patients with advanced melanoma, focusing firstly on patients with stage IV melanoma. We also review treatments available to reduce the risk of a melanoma returning – these treatments can be given either before (“neoadjuvantly”) or after (“adjuvantly”) a melanoma is surgically removed, but only the latter is currently approved.

CLASSES OF THERAPEUTIC AGENTS
Immunotherapy

Immune checkpoint inhibitors (CPIs) are a form of immunotherapy designed to target key regulators of the immune system. Immune checkpoints provide stimulatory or inhibitory control of immunity. Tumours can use the inhibitory pathways to protect themselves from being targeted by the immune system. CPIs currently in clinical use act to block these negative pathways enabling T-cells to recognise cancer cells more efficiently. Agonists for stimulatory pathways are also in clinical development. CPIs were the first class of therapy shown to improve the overall survival (OS) for patients with advanced melanoma and provide hope of durable, long-term responses in a subset of patients. The most extensively studied immune checkpoint receptors are cytotoxic T-lymphocyte-associated protein-4 (CTLA-4) and programmed cell death protein-1 (PD-1). CTLA-4 and PD-1 induce T-cell suppression through non-overlapping mechanisms and likely impact different populations of T-cells during different phases of the immune response (CTLA-4 during priming and PD-1 during the effector phase), providing a mechanistic rationale for the combination of CTLA-4 and PD-1 blockade. CTLA-4. Based on promising antitumour activity in preclinical cancer models (2), CTLA-4-blocking antibodies have been developed. Ipilimumab is a fully human monoclonal antibody of the IgG1 isotype that...
inhibits CTLA-4 leading to enhanced T-cell activation. For T-cell activation to occur, two sequential signals are required (3–5). Firstly, antigens presented in context with the major histocompatibility complex (MHC) I or II on specialised antigen-presenting cells (APCs) must bind with T-cell receptors (TCRs). Following this, there is a translation of TCR stimulation into T-cell activation which requires a costimulatory signal, occurring when B7 surface molecules on the APC bind with CD28 T-cell-surface receptors. Subsequently, T-cell surface expression of CTLA-4 occurs, competitively inhibiting the binding of B7 to CD28, preventing the costimulatory signal and dampening down T-cell activation and proliferation. Treatment can be associated with mechanism-based, immune-related adverse events more frequently than anti-PD-1 treatment.

A second CTLA-4-blocking antibody, tremelimumab, has been developed. Tremelimumab is a fully human anti-CTLA-4 monoclonal antibody of the IgG2 isotype. However, tremelimumab failed to reach its primary endpoint of improved OS compared to standard-of-care chemotherapy for patients with previously untreated, unresectable stage III or IV melanoma (6). Clinical development of tremelimumab is ongoing in a number of non-melanoma cancers.

\[ PD-1 \text{ } \]
Like CTLA-4, PD-1 inhibits T-cell activity and is expressed by activated T-cells. However, instead of competitively inhibiting co-stimulation by interfering with CD28/B7 ligand interaction, PD-1 negatively regulates TCR-signalling events. While CTLA-4 inhibits T-cells during the priming phase of immune responses, PD-1 is thought to inhibit activated T-cells at a later stage in peripheral tissues, playing a critical role in the maintenance of peripheral T-cell tolerance.

The first anti-PD-1 blocking antibody developed was nivolumab, a fully human monoclonal antibody of the IgG4 isotype that binds to PD-1, preventing it from interacting with its ligands. Pembrolizumab was the second anti-PD-1 blocking antibody to be used in advanced melanoma; like nivolumab, pembrolizumab is a fully human monoclonal antibody of the IgG4 isotype that binds to human PD-1 preventing ligand interaction. Nivolumab and pembrolizumab are clinically comparable in terms of efficacy and toxicity as monotherapy for inoperable melanoma (despite the absence of any head-to-head comparison), but only nivolumab is licensed for delivery as a combination with ipilimumab. The subtle preclinical and molecular differences between these two agents have been described by Fessas et al. (7). Compared with ipilimumab, anti-PD-1 blockade with pembrolizumab has been shown to have a superior clinical efficacy and improved toxicity profile with fewer SAEs and fewer patients requiring early treatment withdrawal (8).

\[ \text{Oncolytic virus therapy.} \]
Oncolytic viruses are a novel class of intratumoural immunotherapies that show promise for treating solid tumours. Talimogene laherparvec (T-VEC) is a first-in-class, genetically modified, herpes simplex virus type 1-based oncolytic immunotherapy approved for the local treatment of unresectable cutaneous, subcutaneous and nodal lesions in patients with melanoma recurrent after initial surgery. The mechanism of action and clinical applications of T-VEC are described in detail by Raman et al. (9). The key study to note in the context of advanced melanoma is the OPTiM study which randomised 436 participants in a 2:1 ratio to receive intratumoural T-VEC or subcutaneous re-combinant granulocyte macrophage colony-stimulating factor (GM-CSF). OPTiM first reported positive findings in late 2015 (10), and recently published final analyses confirmed T-VEC’s association with durable complete responses that were associated with prolonged survival (11).

\[ \text{Targeted therapy.} \]
The vast majority of cutaneous melanomas harbour mutations in genes of key signalling pathways. Yet, only a small number of these are considered to be driver mutations due to their active role(s) in cancer development and progression; the others are seen as coincidental passenger mutations that are dispensable for cancer cell viability and develop over the course of tumour evolution (12, 13). The mitogen-activated protein kinase (MAPK) pathway is a complex cascade requiring sequential phosphorylation of the different pathway components. In normal cells, when MAPK activation occurs, it leads to cell growth and differentiation. In cells harbouring BRAF\text{V600E} mutations, the normal process of negative feedback does not occur and this results in permanent MAPK pathway activation, leading to uncontrolled proliferation. This pathway offers various points at which the protein cascade can be blocked. Mutant BRAF is a “driver oncogene” as mutant BRAF inactivation can induce cancer cell toxicity due to an acquired dependency of cancer cells on oncogenic, mutant forms of BRAF (14). Targeted inactivation of BRAF by pharmacologic inhibitors is an archetypal example of targeted therapy in cancer (14, 15). The recognition of key molecular mutation, BRAF\text{V600E} mutation, provided new therapeutic opportunities and facilitated the development of promising small molecule inhibitory compounds later on. Approximately 40% of melanomas harbour a BRAF mutation (16, 17), the most common being BRAF\text{V600E}, followed by BRAF\text{V600K} and rarer genotypes (18).

MEK is the next kinase down from BRAF on the MAPK cascade. BRAF inhibition is the most established form of targeted therapy in melanoma and produces rapid, but often short-lived, tumour regression in the majority of patients. When MEK inhibition is added to BRAF inhibition, increased efficacy and reduced toxicity are seen. Indeed, the combination of BRAF and MEK inhibition offer greater inhibition of MAPK signalling and result in longer durations of response, higher rates
of tumour response, and less cutaneous toxicity often observed from paradoxical MAPK pathway activation with BRAF inhibitor monotherapy (19). The development of acquired resistance to combination BRAF and MEK inhibitor therapy, along with tumour heterogeneity, are formidable obstacles in the treatment of patients with advanced melanoma.

**BRAF inhibitors.** The first BRAF inhibiting tyrosine kinase inhibitor (TKI) approved by the US Food and Drug Administration (FDA) for melanoma treatment was vemurafenib in 2011 (20). The success of vemurafenib in phase I and II settings (21, 22) and then in the BRIM-3 study (23) encouraged intensive investigation of the molecular mechanisms of pathogenesis in melanoma and development of new therapeutic strategies targeting specific molecules in the MAPK pathway. Dabrafenib followed vemurafenib and is another small molecule agent inhibiting BRAFV600 mutation-positive melanoma cell growth, demonstrating efficacy as a monotherapy in the BREAK-3 study (24). Encorafenib is a second-generation BRAF inhibitor, characterised by a substantially prolonged dissociation half-life (25), and in the phase III COLUMBUS trial demonstrated superior efficacy over vemurafenib monotherapy (26).

**MEK inhibitors.** Preclinical and early studies demonstrated that the addition of a MEK inhibitor to a BRAF inhibitor decreased tumour growth, delaying the development of resistance and reducing occurrence of skin lesions in metastatic melanoma (27). As a result, there has been considerable interest in various combinations of BRAF and MEK inhibition. Trametinib was the first MEK inhibitor approved for the treatment of BRAF-mutated metastatic melanoma naïve to BRAF-inhibition. Trametinib is approved for use in combination with dabrafenib showing efficacy both as a monotherapy when compared to investigator’s choice chemotherapy (28), and when combined with dabrafenib (29, 30). Cobimetinib is another MEK inhibitor which demonstrated efficacy while used in combination with vemurafenib in the CoBRIM study (31), while bimetinib is the most recently-introduced of the MEK inhibitors and has demonstrated efficacy in the COLUMBUS study (26).

**Chemotherapy**

Prior to recent advances, chemotherapy was the backbone of treatment for metastatic melanoma. Studies reported responses in 10–15% of patients with 5 year survival in only 2–6% of patients (32). Despite the poor survival statistics, agents such as dacarbazine or the combination of a platinum agent and a taxol were the standard of care for many years, due to a paucity of other useful therapeutic options. Currently chemotherapy is used infrequently, and primarily when immunotherapy and targeted therapy options have either failed or cannot be used.

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**TREATMENT OF UNRESECTABLE STAGE III-IV AND RECURRENT MELANOMA**

Systemic therapy is indicated for patients with stage III–IV melanoma in whom surgical metastasectomy is not appropriate. Patients with oligometastatic disease should be evaluated for possible metastasectomy, as complete resection of metastatic disease can achieve cure (33, 34). In such cases, adjuvant therapy would then be recommended following complete resection to reduce recurrence risk (discussed later). This section will focus on systemic therapy for inoperable melanoma.

The primary systemic therapy options for patients with metastatic melanoma are CPIs, and, where a BRAF mutation is the driver mutation, MAPK targeted therapies. The presence or absence of a BRAF mutation is currently the only reliable predictive biomarker that can influence the treatment of advanced melanoma and must promptly and accurately be determined. Many different methods for BRAF testing are currently in use internationally (35–37), but a discussion of these is beyond the scope of this review. Targeted MAPK therapy is not indicated in patients without a characteristic BRAF mutation and may indeed be harmful to this patient group.

Whether patients with BRAFV600 mutant melanoma should receive CPIs or MAPK targeted therapy as first line therapy is not always straightforward and prospective head-to-head comparative trials of MAPK inhibitors and CPIs are lacking. A 2019 update of survival in metastatic melanoma reported exploratory analysis of survival data from selected CPI and TKI clinical trials (38). In first line therapy, mean 3-year OS proportions were 41.3% for BRAF plus MEK inhibition, 49.9% for PD-1 inhibition and 58.4% for CTLA-4 plus PD-1 inhibition. Comparison of the mean progression free survival (PFS) and OS curves of kinase inhibition and checkpoint blockade revealed a superiority of combined BRAF plus MEK inhibition within the first 12 months, later changing to a superiority of PD-1 blockers alone or in combination with CTLA-4 blockade. In second-line or higher, BRAF plus MEK inhibition was superior to anti-PD-1 monotherapy throughout the first 3 years; mean 3-year OS proportions were 42.4% for BRAF plus MEK inhibition, and 40.1% for PD-1 inhibition.

**Checkpoint inhibitors**

Table I outlines key phase III CPI studies in melanoma. Nivolumab (39) and pembrolizumab (40, 41) have been established as preferred monotherapy options for inoperable melanoma given their efficacy over standard of care chemotherapies and acceptable toxicity profiles. Checkmate-067 compared nivolumab and ipilimumab as a combination with nivolumab and ipilimumab monotherapies, recently demonstrating an OS of 52% for the combination group at 5 years. This exceptional survival was associated with 59% of patients receiving the combination suffering
grade 3 or 4 adverse events (42). As such, combination PD-1 and CTLA-4 blockade is usually considered only for those patients with a very good performance status, with some institutions and oncologists preferring CPI monotherapy for metastatic disease. Untreated brain metastases represent one particular clinical scenario in which combination CPI offers particular advantage and may be preferred in this instance (43).

**MAPK pathway inhibition**

Overall response rates to vemurafenib, dabrafenib and encorafenib monotherapies are 45%, 51% and 60%, respectively (29, 44, 45). A number of studies have presented clear evidence that the combination of these agents with a MEK inhibitor provide increased efficacy with a reduction in toxicity (Table II). In the COLUMBUS study, encorafenib plus binimetinib showed favourable efficacy compared with encorafenib or vemurafenib monotherapy, with the combination associated with an improved tolerability profile compared with either monotherapies (26). The CoBRIM study showed improved survival of vemurafenib and cobimetinib compared with vemurafenib alone, with no significant difference in toxicity (31). Robert et al. recently analysed pooled extended survival data from COMBI-d and COMBI-v trials (n = 563) which compared dabrafenib and trametinib with dabrafenib and vemurafenib monotherapies, respectively, reporting complete responses in 19% of patients and improved long-term outcomes, with OS rates of 71% and less toxicity seen with the combination of BRAF and MEK inhibition (29).

**Checkpoint and MAPK inhibition combinations**

Increasing evidence suggests that BRAF and MEK inhibition has an immune-modulating effect, enhancing anti-tumour immunity (47–49). Early evidence from treatment of advanced melanoma with BRAF inhibition demonstrated increased expression of PD-1 and its

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**Table I. Landmark checkpoint inhibitor (CPI) trials in metastatic melanoma**

| Trial | Regimen | Patients n | Outcome | G3/4 AEs: |
|---|---|---|---|---|
| Checkmate 066 (40) | Vemurafenib 960 mg BD vs. DTIC 1,000 mg/m² q3w | 418 | 3 years OS: 51.2% vs 21.6% mOS: 37.5 vs 11.2 months | 11.7% vs 17.6% |
| Nivo 1st line | Nivo 3 mg/kg q2w vs. DTIC 1,000 mg/m² q2w | 405 | ORR: 27% vs 10% mOS: 16 vs 14 mo mPFS: 3.1 vs 3.7 mo | 14% vs 34% |
| Checkmate 037 (41) | Nivo 3mg/kg q2w vs. ICC | 295 | PFS at 60 months: 36%* G3/4 AEs: 29%* (Nivo) vs 26%* (ICC) | 59% (Ipi+Nivo) vs 23% (Nivo) vs 28% (Ipi) |
| Nivo 2nd line | Nivo 3 mg/kg q2w vs. ICC | 834 | mOS at 60 months: 32.7% vs 15.9% mPFS at 60 months: 8.4 months vs 3.4 months | 17% vs 50% |
| Checkmate 067 (42) | Comparison of 3 x 3-weekly regimens: Nivo 1mg/kg + Ipi 3 mg/kg q3w vs. Nivo 3 mg/kg q2w vs. Ipi 3 mg/kg x 4 doses | 945 | PFS at 28 months: 36% (pembro 2 mg) vs 38% (pembro 10 mg) vs 30% (ICC) | 13.5% (pembro 2 mg) vs 16.8% (pembro 10 mg) vs 26.3% (ICC) |
| Keynote-006 (8) | Pembrolizumab 10 mg/kg | 405 | ORR: 27% vs 10% mOS: 37.5 vs 11.2 months | 11% vs 17.6% |
| Pembrolizumab 2nd line (Ipi refractory) | Pembrolizumab 2 mg/kg q3w vs. Pembrolizumab 10 mg/kg vs. Pembrolizumab 2nd line (Ipi refractory) | 180 | mOS at 28 months: 13.4 (pembro 2 mg) vs 14.7 (pembro 10 mg) vs 11.0 months | 11% vs 17.6% |

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**Table II. Landmark mitogen-activated protein kinase (MAPK) targeted therapy trials in metastatic melanoma**

| Trial | Regimen | Patients n | Outcome | Toxicity |
|---|---|---|---|---|
| BRIM-3 (23) | Vemurafenib 960 mg BD vs. DTIC 1,000 mg/m² q3w | 675 | mOS: 13.6 vs 9.7 months mPFS: 6.9 vs 1.6 months | Modification/Interruption: 38% vs 16% |
| BREAK-3 (24) | Dabrafenib vs. DTIC | 250 | mPFS: 5.1 months vs 2.7 months | G3/4 AEs: 12.8% vs 17.4% |
| METRIC (28) | Trametinib 2 mg/day vs. ICC | 322 | mPFS: 4.9 vs 1.5 months | G3/4 AEs: 29% vs 12% |
| CoBRIM (31) | Vemurafenib + Cobimetinib 60 mg OD vs. Vemurafenib 960 mg BD + placebo | 495 | 5 year OS: 32% vs 17% | 60% vs 52% |
| COMBI-d (46) | Dabrafenib 150 mg BD + Trametinib 2 mg OD vs. Dabrafenib 150 mg + placebo | 423 | 3 years OS: 44% vs 32% mPFS: 11.0 vs 8.8 months | 48% vs 50% |
| COMBI-v (39) | Dabrafenib 150 mg BD + Trametinib 2 mg OD vs. Vemurafenib 960 mg BD | 704 | 5 years pooled results with COMBI-d: CR in 19%; OS rates of 71% (29) | G3/4 AEs: 58% vs 66% |
| COLUMBUS (26) | Encorafenib 450 mg OD + Binimetinib 45 mg BD (Combo) vs. Encorafenib 300 mg OD vs. Vemurafenib 960 mg BD | 577 | mOS: 33.6% (combo) vs 23.5 months (enco) vs 16.9% (vem) mPFS: 14.9 months (combo) vs 9.6 months (enco) vs 7.3 months (vem) | 68% (combo) vs 66% (vem) |

AEs: adverse events; OD: once daily; BD: twice daily; mOS: median overall survival; HR: hazard ratio; mPFS: median progression-free survival; PD: progressive disease; G: grade; AE: adverse event; TRAE: treatment-related adverse event; Ipi: Ipilimumab; Nivo: Nivolumab; Pembro: Pembrolizumab; DTIC: Dacarbazine; ICC: investigator’s choice chemotherapy.

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Theme issue: Skin malignancies
The role of adjuvant therapy in patients with resected stage III melanoma is a rapidly evolving field. Interferon was the first agent shown to have utility in this space, however, advances in both targeted therapies and immunotherapies have led to a number of practice-changing adjuvant trials in resected stage III and IV disease. By eliminating the micrometastatic disease that remains after surgery, adjuvant systemic therapy aims to reduce disease recurrence and ultimately improve rates of cure following surgical resection of locoregional or stage IV disease. Patients with resected stage III or IV disease have significant differences in predicted survival at 5 years ranging from approximately 80% for stage IIIa disease to less than 20% for resected stage IIIId disease (58). Adjuvant treatment with either CPI or MAPK targeted therapy have dramatically changed outcomes for this patient group, with approximately 50% increased recurrence-free survival (RFS) for both treatment approaches (59–62). CPIs and MAPK targeted therapies have not been directly compared in phase III studies and there is currently no clear consensus on choice of approach for patients with a BRAFV600E mutation in the adjuvant setting.

For patients with stage I and II primary tumours and a negative sentinel lymph node biopsy, there is presently no indication for adjuvant therapy (63). It is worth noting that patients with high risk (primary tumour >4 mm, or >2 mm with ulceration) but node negative tumours were excluded from the phase III clinical trials that evaluated nivolumab, ipilimumab and the targeted therapy doublet of dabrafenib and trametinib (62, 64, 65). As such, data on adjuvant therapy in this cohort of patients is not available and is currently under investigation.

**Adjuvant checkpoint inhibitors**

As already discussed, CPI represents an important advance in the treatment of patients with inoperable melanoma. These results led to the evaluation of these agents in the adjuvant setting for patients at high risk of recurrence following initial surgery. Adjuvant treatment with ipilimumab at 10 mg/kg dosing was shown to have a 10% absolute improvement in OS and RFS, but toxicity and high treatment-related death rates limited its widespread use and it was never licensed for this indication in Europe (66). Only 13.4% of patients completed the full planned course of treatment, and nearly 40% of patients discontinued treatment after the first 4 doses due to treatment-related side effects. Adjuvant anti-PD-1 therapy has been tested in two large phase III studies, Checkmate 238 and Keynote 054, which have established nivolumab and pembrolizumab as the CPIs of choice for the adjuvant treatment of resected melanoma (60, 67). Table IV summarises the key trials in this setting.

**Adjuvant targeted therapy**

A key study in this context is COMBI-AD, a study of 870 Stage III BRAF mutant melanoma patients in the adjuvant setting following excision and lymphadenectomy (61, 64). They were randomised to the combination arm of dabrafenib and trametinib, or to matching placebos

### Table III. Landmark check-point/tyrosine kinase inhibitor (CPI-TKI) targeted therapy trials in metastatic melanoma

| Trial            | Regimen                                                                 | Patients n | Outcome | Toxicity |
|------------------|-------------------------------------------------------------------------|------------|---------|----------|
| Keynote 022     | Pembrolizumab 2 mg/kg + Dabrafenib 150 mg BD + Tremetibib 2 mg          | 120        | mPFS: 16.0 vs 10.3 mDOR: 18.7 months vs 12.5 mOS: NR vs 23.4 | G3-5 AEs: 70% vs 45% |
| NCT02130466      | pembrolizumab 2 mg OD vs Placebo + Dabrafenib 150 mg BD + Tremetibib 2 mg OD |            |         |          |
| IMspire150       | Atezolizumab 840 mg D1 and D15 + Vemurafenib 960 mg BD + Cobimetinib 60 mg OD | 514        | PFS: 15.1 vs 10.6 months 2 years OS: 60.4% vs 53.1% | G3-5 AEs: 33.5% vs 28.8% |
| NCT02908672      | Atezolizumab 840 mg D1 and D15 + Vemurafenib 960 mg BD + Cobimetinib 60 mg OD |            |         |          |
| COMBI-I          | Spartializumab 400mg q4W + Dabrafenib 150 mg BD + Tremetibib 36          | 12 months PFS: 65.3% 12 months OS: 85.9% | 75% had G3/4 AEs |
| NCT02967692      | Spartializumab 400mg q4W + Dabrafenib 150 mg BD + Tremetibib 36          | 12 months PFS: 65.3% 12 months OS: 85.9% | 75% had G3/4 AEs |

AEs: adverse events; OD: once daily; BD: twice daily; mOS: median overall survival; HR: hazard ratio; mPFS: median progression-free survival; mDOR: median duration of response; ORR: overall response rate; NR: not reached; PD: progressive disease; G: grade; AE: adverse event; DTIC: Dacarbazine; ICC: investigator’s choice chemotherapy.
for one year. The primary endpoint, RFS, was longer with dabrafenib and trametinib than with placebo (4-year rate: 54% vs 38%; hazard ratio [HR] 0.49, 95% CI 0.40–0.59), with treatment benefits observed irrespective of baseline factors, according to subgroup analysis (61). Vemurafenib was compared to placebo in the adjuvant BRIM8 study demonstrating efficacy but high rates of grade 3/4 toxicity (68).

**NEOADJUVANT THERAPY FOR EARLY MELANOMA**

Given the success of immunotherapies and targeted therapies for the treatment of advanced melanoma, the natural extension is to identify the role of these therapies in the neoadjuvant setting, with a wealth of clinical trials currently underway. Patients with clinically detectable stage III melanoma represent a high-risk population with poor outcomes when treated with upfront surgery alone and are obvious candidates for investigation of neoadjuvant therapy. However, the clear need to carefully evaluate short-term clinical endpoints such as RFS, and long-term endpoints of neoadjuvant therapy against those of adjuvant therapy remains. Neoadjuvant therapy for melanoma is not presently standard-of-care but represents an active area of research with a large number of completed and recruiting trials with differing designs, endpoints, and methods of analysis under investigation. **Table V** illustrates those neoadjuvant (preoperative therapy) trials which have reported data.

One study of note is OPACIN-NEO study which reported in 2018 (69). OPACIN-NEO examined neoadjuvant combination CPI with 3 different regimens of ipilimumab and nivolumab. A combination of ipilimumab at 1 mg/kg combined with nivolumab at 3 mg/kg given 3-weekly for two cycles was chosen to take forward into later phase studies, as this combination had a response rate of 77%, with responders experiencing excellent outcomes to date. If more mature data confirm these early observations, this schedule will be tested in randomised phase 3 studies versus adjuvant therapies, which are the current standard-of-care systemic therapy for patients with stage III melanoma.

**FUTURE DIRECTIONS AND CONCLUSION**

The investigation of new immunotherapy and/or targeted therapy combinations, such as anti-PD-1/anti-CTLA-4 CPIs with other immunotherapies (e.g. indoleamine 2,3 dioxygenase inhibitors, antilymphocyte receptor, pegylated interleukin-2), combination targeted therapies (e.g. MEK and CDK4/6 co-inhibition), and the combined use of immunotherapy and continued research on targeted therapy (e.g. the triplet combination of BRAF/MEK inhibition with anti–PD-1s) are keys for the future of systemic therapy for advanced melanoma. The identification of novel therapeutic targets in the MAPK pathway provides opportunity to improve outcomes by overcoming de novo and acquired resistance to BRAF/MEK inhibition. Adoptive cell transfer may have a potential role in patients whose disease has progressed following CPI. Altogether, these new approaches offer potential to build upon past advances and improve long-term survival outcomes for patients with melanoma.

This decade has brought significantly improved outcomes for patients with advanced melanoma with the advent of immunotherapies and targeted treatments that have utility in a variety of settings. However, responses to treatment are heterogeneous and not always durable. Further advances are required, and several emerging strategies are of particular interest.

**Conflicts of interest:** KAL has no conflicts of interest to report. PN reports personal fees from AstraZeneca, BMS, Merck, Immunocore, Pfizer, Ipsen, 4SC, Pierre Fabre and Roche.
Table V. Neoadjuvant trials with available data

| Trial | Eligible patients n | Regimen | Median follow-up (months) | Results | TRAEs |
|-------|---------------------|---------|--------------------------|---------|-------|
| **IMMUNOTHERAPY** | | | | | |
| NCT00972933 (70) | Clinical stage IIIB or IIIC and oligometastatic stage IV; n = 35 | Two neoadjuvant doses of ipi (10 mg/kg), surgery, followed by two adjuvant doses of ipi | 18 | RFS: 11 months | No pPR or pCR reported | G3 AEs: 32% |
| NCT02437279 (71) | Clinical stage III; 10 per group | Surgery plus 12-week adjuvant ipi (3 mg/kg) and nivo (1 mg/kg); 6 weeks of neoadjuvant and 6 weeks of adjuvant ipi (3 mg/kg) and nivo (1 mg/kg) | 32 | 30% pCR, 40% near pCR, 0% pPR | G4/3 adverse events: 90% of participants in the surgery group vs 90% of participant in the neoadjuvant therapy group |
| NCT02519322 (72) | Clinical stage III and oligometastatic stage IV; 12 participants in the nivo-only group and 11 in the ipi plus nivo group | 4 doses of nivo (3 mg/kg) neoadjuvant therapy, surgery, and 24 weeks of nivo adjuvant therapy; 3 courses of ipi (3 mg/kg) plus nivo (1 mg/kg) neoadjuvant therapy, surgery, and 24 weeks of adjuvant nivo | 20 | Group A: pCR 45% Group B: pCR 25% | Nivolumab-only: 8% participants had G3 AEs; ipi plus nivo: 73% participants had G3 AEs; No G4/5 AEs in any group |
| NCT02977052 (69) | OpACIN-neo 2019 | Group A: two courses of ipi (3 mg/kg) plus nivo (1 mg/kg) once every 3 weeks; Group B: two courses of ipi (1 mg/kg) plus nivo (3 mg/kg) every 3 weeks; Group C: two courses of ipi (3 mg/kg) once every 3 weeks plus two courses of nivo (1 mg/kg) once every 2 weeks | 8.3 | 43% of non-pCRs relapsed; no relapses reported in the other response groups | G3/4 AEs: 40% in group A vs 20% in group B vs 50% in group C |
| NCT01608594 (73) | Clinically detectable locally and/or regionally advanced melanoma; n = 28 | Ipilimumab 3 or 10 mg/kg high-dose interferon | 32 | 32% pCR | At median follow-up of 32 months, 10/11 patients with either pCR or minimal residual disease remained disease free |
| NCT02339324 (74) | Stage 3 and 4 resected (5 x IIIB, 11 x IIIC and 4 x IV); n = 20 | Pembrolizumab 200 mg with high-dose interferon | 11 | 35% pCR | 90% of patients had to stop early due to G4/4 toxicities |
| **TARGETED THERAPY** | | | | | |
| NCT02211775 (75) | Clinical stage IIIB or IIIC and oligometastatic stage IV with BRAFV600E/V600K mutation; n = 21 | Neoadjuvant dabrafenib (150 mg twice a day) plus trametinib (2 mg daily) for 8 weeks followed by surgery and 44 weeks of the same adjuvant treatment versus surgery | 18.6 | pPR 17% and pCR 58% | A: G3: 47% of participants in the neoadjuvant systemic therapy group had G3 AEs |
| NCT01972347 (76) | NeoCombi 2019 | Dabrafenib (150 mg twice a day) plus trametinib (2 mg daily); 12 weeks neoadjuvant therapy and 40 weeks of adjuvant therapy | 27 | 23 mo of overall RFS (30 mo of pCR, 18 mo of non-pCR) | 57% participant had any grade 3 adverse events; 3% had any G4 AEs and 26% had surgical G3 AEs; 26% had drug-related grade 3 and 4 AEs |
| Slott et al. (77) | Stage III Of 15, 6 underwent surgery | Vemurafenib 960 mg BID or Dabrafenib 25.4 | | pPR 33% and pCR 33% | Dose reduction or discontinuation because of toxicities occurred in 10/15 patients |
| Zippel et al. (78) | Stage III | Vemurafenib 960 mg BID or Dabrafenib 20 | | pPR 62% and pCR 31% | N/a |
| Ergul et al. (79) | Stage IIIIC and IV | Vemurafenib Dabrafenib + Trametinib | 25 | pCR 35% | Not reported |

pCR: pathological complete response; mo: months; G: grade; TRAE: treatment related adverse events; AE: adverse event; ipi: ipilimumab; nivo: nivolumab; pPR: pathological partial response; pCR: pathological complete response.

REFERENCES

1. Domingues B, Lopes JM, Soares P, Pópulo H. Melanoma treatment in review. Immunotargets Ther 2018; 7: 35–49.
2. Keler T, Halk E, Vitale L, O’Neill T, Blanset D, Lee S, et al. Activity and safety of CTLA-4 blockade combined with vaccines in cynomolgus macaques. J Immunol 2003; 171: 6251–6259.
3. Boasberg P, Hamid O, Day S. Ipilimumab: unleashing the power of the immune system through CTLA-4 blockade. Semin Oncol 2010; 37: 440–449.
4. Hoos A, Ibrahim R, Korman A, Abdallah K, Berman D, Shahabi V, et al. Development of ipilimumab: contribution to a new paradigm for cancer immunotherapy. Semin Oncol 2010; 37: 533–546.
5. Weber J. Immune checkpoint proteins: a new therapeutic paradigm for cancer – preclinical background: CTLA-4 and PD-1 blockade. Semin Oncol 2010; 37: 430–439.
6. Ribas A, Kefford R, Marshall MA, Punt CIA, Haanen JB, Marmol M, et al. Phase III randomized clinical trial comparing tremelimumab with standard-of-care chemotherapy in patients with advanced melanoma. J Clin Oncol 2013; 31: 616–622.
7. Fessas P, Lee H, Ikemizu S, Janowitz T. A molecular and preclinical comparison of the PD-1-targeted T-cell checkpoint inhibitors nivolumab and pembrolizumab. Semin Oncol 2017; 44: 136–140.
8. Robert C, Ribas A, Schachter J, Arance A, Grob J-J, Mortier L, et al. Pembrolizumab versus ipilimumab in advanced melanoma (KEYNOTE-006): post-hoc 5-year results from an open-label, multicentre, randomised, controlled, phase 3 study. Lancet Oncol 2019; 20: 1239–1251.
9. Raman SS, Hecht JR, Chan E. Talimogene laherparepvec: durable response rate in patients with advanced melanoma. J Clin Oncol 2013; 31: 616–622.
10. Andtbacka RH, Kaufman HL, Collichio F, Amatruda T, Senzer N, Chesney J, et al. Talimogene laherparepvec improves overall survival in patients with advanced melanoma treated with ipilimumab. J Clin Oncol 2015; 33: 2780–2788.
11. Andtbacka RHI, Collichio F, Harrington KJ, Middleton MR,
Downey G, Öhring K, et al. Final analyses of OPTIM: a randomized phase III trial of talimogene laherparepvec versus granulocyte-macrophage colony-stimulating factor in unresectable stage III–IV melanoma. J ImmunoTher Cancer 2019; 7: 145.

12. Clarke CN, Katsonis P, Hsu TK, Koire AM, Silva-Figueroa A, Christakis I, et al. Comprehensive genomic characterization of parathyroid cancer identifies novel candidate driver mutations and core pathways. J Endocr Soc 2019; 3: 544–559.

13. Kone HS, Minna JD, White MA, GNAS meets TCGA to illuminate mechanisms of cancer predisposition. Cell 2013; 152: 387–389.

14. Rustgi AK. BRAF: a driver of the serrated pathway in colon cancer. Cancer Cell 2013; 24: 1–2.

15. Hofer S, Berthod G, Riklin C, Rushing E, Feilchenfeld J. BRAF V600E mutation: A treatable driver mutation in pleomorphic xanthoastrocytoma (PXA). Acta Oncol 2016; 55: 122–123.

16. Colombino M, Capone M, Lissia A, Assoua S, Rubino C, De Giorgi V, et al. BRAF/NRAS mutation frequencies among primary tumors and metastases in patients with melanoma. J Clin Oncol 2012; 30: 2522–2529.

17. Curtin JA, Fridlyand J, Kageshita T, Patel HN, Busam KJ, Kutzner H, et al. Distinct sets of genetic alterations in melanoma. N Engl J Med 2005; 353: 2135–2147.

18. Menzies AM, Haydu LE, Visintin L, Carlino MS, Howle JR, Thompson JF, et al. Distinguishing clinicopathologic features of patients with V600E and V600K BRAF-mutant metastatic melanoma. Clin Cancer Res 2012; 18: 3242–3249.

19. Erogol Z, Ribas A, Comin-Anduix B, Dummer R, and MEK inhibitors for melanoma: latest evidence and place in therapy. Ther Adv Med Oncol 2016; 8: 48–56.

20. Chapman PB, Hauschild A, Robert C, Haanen JB, Ascierto P, Larkin J, et al. Improved survival with vemurafenib in melanoma with BRAF V600E mutation. N Engl J Med 2011; 364: 2507–2516.

21. Flaherty KT, Puzanov I, Kim KB, Ribas A, McArthur GA, Sosman JA, et al. Inhibition of Mutated, Activated BRAF in Metastatic Melanoma. N Engl J Med 2010; 363: 809-819.

22. Ribas A, Kim KB, Schuchter LM, Gonzalez R, Pavlick AC, Weber JS, et al. BRIM-2: An open-label, multicenter phase II study of vemurafenib in previously treated patients with BRAF V600E mutation-positive metastatic melanoma. J Clin Oncol 2011; 29: 8509.

23. Chapman PB, Robert C, Larkin J, Haanen JB, Ribas A, Hogg D, et al. Vemurafenib in patients with BRAFV600 mutation-positive metastatic melanoma: final overall survival results of the randomized BRIM-3 study. Annals Oncol 2017; 28: 2581–2587.

24. Hauschild A, Grob JJ, Demidov LV, Jouary T, Gutzmer R, Hauschild A, et al. Combined BRAF and MEK inhibition versus vemurafenib or encorafenib in patients with BRAF V600-mutant melanoma. Eur J Cancer 2020; 126: 33–44.

25. Kim KB, Keffer F, Pavlick AC, Infante JR, Ribas A, Sosman JA, et al. Phase II study of the MEK1/MEK2 inhibitor Trametinib in patients with metastatic BRAF-mutant cutaneous melanoma previously treated with or without a BRAF inhibitor. J Clin Oncol 2013; 31: 482–489.

26. Robert C, Flaherty K, Nathan P, Hersey P, Garbe C, Milhem M, et al. Five-year outcomes from a phase 3 METRIC study in patients with BRAF V600 E/K-mutant advanced or metastatic melanoma. Eur J Cancer 2020; 126: 33–44.

27. Kim KB, Keffer F, Pavlick AC, Infante JR, Ribas A, Sosman JA, et al. Phase II study of the MEK1/MEK2 inhibitor Trametinib in patients with metastatic BRAF-mutant cutaneous melanoma previously treated with or without a BRAF inhibitor. J Clin Oncol 2013; 31: 482–489.

28. Robert C, Flaherty K, Nathan P, Hersey P, Garbe C, Milhem M, et al. Five-year outcomes from a phase 3 METRIC study in patients with BRAF V600 E/K-mutant advanced or metastatic melanoma. Eur J Cancer 2019; 109: 61–69.

29. Long GV, Stroyakovsky D, Dogramaci L, Shah A, Klaasa R, Savage KJ. Long-term survival in patients with metastatic melanoma treated with DTIC or temozolomide. Oncologist 2010; 15: 765–771.

30. Leung AM, Hari DM, Morton DL. Surgery for distant melanoma metastasis. Cancer J 2012; 18: 176–184.

31. Ollila DW, Hsueh EC, Stern SL, Morton DL. Metastasectomy for recurrent stage IV melanoma. J Surg Oncol 1999: 71; 209–213.

32. Finn RS, Dering J, Konklin D, Kalous O, Cohen DJ, Desai AJ, et al. PD 0329911, a selective cyclin D kinase 4/6 inhibitor, preferentially inhibits proliferation of luminal estrogen receptor-positive human breast cancer cell lines in vitro. Breast Cancer Res 2009; 11: R77.

33. Finn RS, Crown JP, Lang I, Boer K, Bondarenko IM, Kulyk SO, et al. The cyclin-dependent kinase 4/6 inhibitor palbociclib in combination with letrozole versus letrozole alone as first-line treatment of oestrogen receptor-positive, HER2-negative, advanced breast cancer (PALOMA-1/TRIO-18): a randomised phase 2 study. Lancet Oncol 2015; 16: 25–35.

34. Ihle MA, Fassunke C, Bommarius V, Grunewald I, Schlaak M, Kreuzberg N, et al. Comparison of high resolution melting analysis, pyrosequencing, next generation sequencing analysis and immunohistochemistry to conventional Sanger sequencing for the detection of p.V600E and non-p.V600E BRAF mutations. BMC Cancer 2014; 14: 13.

35. Uqviel S, Rohmel J, Ascierto PA, Becker JC, Flaherty KT, Grob JJ, et al. Survival of patients with advanced metastatic melanoma: The impact of MAP kinase pathway inhibition and immune checkpoint inhibition - Update 2019. Eur J Cancer 2020; 130: 126-138.

36. Hamid O, Puzanov I, Dunhmer R, Schachter J, Daud A, Schadendorf D, et al. Final analysis of a randomised trial comparing pembrolizumab versus investigator-choice chemotherapy for ipilimumab-refractory advanced melanoma. Eur J Cancer 2017; 86: 37–45.

37. Ascierto PA, Long GV, Robert C, Bradley B, Dutriaux C, Di Giacomo AM, et al. Survival outcomes in patients with previously untreated advanced melanoma treated with nivolumab plus ipilimumab therapy: three-year follow-up of a randomized phase 3 trial. JAMA Oncol 2019; 5: 187–194.

38. Larkin J, Minor D, D’Angelo S, Nneys B, Smylie M, Miller WH, Jr., et al. Overall survival in patients with advanced melanoma who received nivolumab plus investigator’s choice chemotherapy in checkmate 037: a randomized, controlled, open-label phase III trial. J Clin Oncol 2018; 36: 383–390.

39. Larkin J, Chiorion-Silveni V, Gonzalez R, Grob J-R, Rutkowski P, Lao CD, et al. Five-year survival with combined nivolumab and ipilimumab in advanced melanoma. N Engl J Med 2019; 381: 1535–1546.

40. Tawbi HA, Forsyth PA, Alagzi A, Hamid O, Hodi FS, Moschos SJ, et al. Combined Nivolumab and Ipilimumab in Melanoma Metastatic to the Brain. N Engl J Med 2018; 379: 722–730.

41. Delord JP, Robert C, Nyakas M, McArthur GA, Kudchakar R, Majeral A, et al. Phase I dose-escalation and -expansion study of the BRAF inhibitor encorafenib (LGX818) in metastatic BRAF-mutant melanoma. Clin Cancer Res 2017; 23: 5337–5348.

42. Larkin J, Ascierto PA, Dreno B, Atkinson V, Lischka G, Maio M, et al. Combined vemurafenib and cobimetinib in BRAF-mutated melanoma. N Engl J Med 2014; 371: 1867–1876.

43. Long GV, Flaherty KT, Stroyakovsky D, Dogramaci L, Heuvenko E, de Braud F, et al. Dabrafenib plus trametinib versus dabrafenib monotherapy in patients with metastatic BRAF V600E/K-mutant melanoma: long-term survival and safety analysis of a phase 3 study. Ann Oncol 2017; 28: 1631–1639.

44. Hu-Lieskovan S, Mok S, Hormet Moreno B, Tsoi J, Robert L, Goedert L, et al. Improved antitumor activity of immu-
notherapy with BRAF and MEK inhibitors in BRAF(V600E) melanoma. Sci Transl Med 2015; 7: 279ra41.

48. Ascierto PA, Dummer R. Immunological effects of BRAF+MEK inhibition. Oncoimmunol 2018; 7: e1468955.

49. Deken MA, Gadiot J, Jordanova ES, Lacroix R, van Gool M, Kroon P, et al. Targeting the MAPK and PI3K pathways in combination with PD1 blockade in melanoma. Oncoimmunol 2016; 5: e1238557.

50. Frederick DT, Piris A, Cogdill AP, Cooper ZA, Lezcano C, Ferrone CR, et al. BRAF inhibition is associated with enhanced melanoma antigen expression and a more favorable tumor microenvironment in patients with metastatic melanoma. Clin Cancer Res 2013; 19: 1225–1231.

51. Ribas A, Hodi FS, Callahan M, Konto C, Wolchok J. Hepatotoxicity with combination of vemurafenib and ipilimumab. N Engl J Med 2013; 368: 1365–1366.

52. Vella LJ, Pasam A, Dimopoulos N, Andrews M, Knights A, Puaux AL, et al. MEK inhibition, alone or in combination with BRAF inhibition, affects multiple functions of isolated normal human lymphocytes and dendritic cells. Cancer Immunol Res 2014; 2: 351–360.

53. Ebert PJR, Cheung J, Yang Y, McNamara E, Hong R, Ross MI, et al. Identification of the optimal combination dosing schedule of neoadjuvant ipilimumab plus nivolumab in macroscopic stage III melanoma (OpACINneo): a multicentre, phase 2, randomised, controlled trial. Lancet Oncol 2019; 20: 910–920.

54. Ascierto PA, Ferrucci PF, Fisher R, Del Vecchio M, Atkinson V, Schmidt H, et al. Dabrafenib, trametinib and pembrolizumab or placebo in BRAF-mutant melanoma. Nature Med 2019; 25: 941–946.

55. Amaria RN, Prieto PA, Tetzlaff MT, Reuben A, Andrews MC, Ross MI, et al. Neoadjuvant immune checkpoint blockade in high-risk resectable melanoma: a single-centre, open-label, single-arm, phase 3 trial. Lancet Oncol 2018; 19: 510–520.

56. Rozeman EA, Menzies AM, van Akkooi ACJ, Adhikari C, Bierman C, van de Wiel BA, et al. Identification of the optimal combination dosing schedule of neoadjuvant ipilimumab plus nivolumab in macroscopic stage III melanoma (OpACINneo): a multicentre, phase 2, randomised, controlled trial. Lancet Oncol 2019; 20: 948–960.

57. Retseck J, Nasr A, Lin Y, Lin H, Mendiratta P, Butterfield LH, et al. Long term impact of CTLA4 blockade immunotherapy on regulatory and effector immune responses in patients with melanoma. J Transl Med 2018; 16: 184.

58. Rozeman EA, Blank CU, Van Akkooi ACJ, Kvistborg P, Fanchi L, Van Thienen IV, et al. Neoadjuvant ipilimumab + nivolumab (IPI+NIVO) in palpable stage III melanoma: Updated data from the OpACIN trial and first immunological analyses. J Clin Oncol 2017; 35: 9586.

59. Amaria RN, Reddy SM, Tawbi HA, Davies MA, Ross MI, Glitza IC, et al. Neoadjuvant immune checkpoint blockade in high-risk resectable melanoma: Evidence-based changes to melanoma staging in the American Joint Committee on Cancer eighth edition cancer staging manual. CA Cancer J Clin 2017; 67: 472–492.

60. Long GV, Saw RPM, Lo S, Nieweg OE, Gonzalez-Moreno C, van de Wiel BA, et al. Identification of the optimal combination dosing schedule of neoadjuvant ipilimumab plus nivolumab in locally/regionally advanced melanoma: high dose IFN-α2b in locally/regionally advanced melanoma: safety, efficacy and impact on T-cell repertoire. J Immunother Cancer 2018; 6: 112.

61. Arance AM, Lin Y, Lin H, Rahman Z, Vallabhaneni P, Mendiratta P, et al. Neoadjuvant ipilimumab (3 mg/kg or 10 mg/kg) and high dose IFN-α2b in locally/regionally advanced melanoma: safety, efficacy and impact on T-cell repertoire. J Immunother Cancer 2018; 6: 112.

62. Tzakis AG, Tzavaras MT, et al. Neoadjuvant combination immunotherapy with pembrolizumab and high dose IFN-α2b in locally/regionally advanced melanoma. J Clin Oncol 2018; 36: 181.

63. Amaria RN, Prieto PA, Tetzlaff MT, Reuben A, Andrews MC, Ross MI, et al. Neoadjuvant plus adjuvant dabrafenib and trametinib versus standard of care in patients with high-risk, surgically resectable melanoma: a single-centre, open-label, randomised, phase 2 trial. Lancet Oncol 2018; 19: 181–193.

64. Long GV, Saw RPM, Lo S, Nieweg OE, Shannon KF, Gonzalez-Moreno C, et al. Neoadjuvant dabrafenib combined with trametinib for resectable, stage IIIIB-C, BRAF(V600) mutation-positive melanoma (NeoCombI): a single-arm, open-label, single-centre, phase 2 trial. Lancet Oncol 2019; 20: 961–971.

65. Slot S, Zager JS, Kudchadkar RR, Messina JL, Benedict JG, Gonzalez-Moreno C, et al. BRAF inhibition for advanced locoregional BRAF V600E mutation-positive melanoma: a potential neoadjuvant strategy. Melanoma Res 2016; 26: 83–87.

66. Zippel D, Markel G, Shapiro-Frommer R, Ben-Betzalel G, Goitzen D, Ben-Ami E, et al. Perioperative BRAF inhibitors in locally advanced stage III melanoma. J Surg Oncol 2017; 116: 856–861.

67. Ergoul Z, Khushalani NI, Rich J, Sarnaik A, Zager JS, Markowitz J, et al. Patterns of histologic response to neoadjuvant targeted therapy in patients with BRAF mutant melanoma. J Clin Oncol 2017; 35: 9584.

68. Hauschild A, Dummer R, Schadendorf D, Santinami M, Atkinson V, Mandala M, et al. Longer follow-up confirms relapse-free survival benefit with adjuvant dabrafenib plus trametinib in patients with resected BRAF V600-mutant stage III melanoma. J Clin Oncol 2018; 36: 3441–3449.

69. Weber JS, Mandala M, Del Vecchio M, Gogas H, Arance AM, Cowey CL, et al. Adjuvant nivolumab versus placebo after complete resection of high-risk stage III melanoma (EORTC 18071): a randomised, double-blind, phase 3 trial. Lancet Oncol 2015; 16: S22–S30.

70. Hauschild A, Dummer R, Wolchok JD, Schmidt H, et al. Adjuvant ipilimumab versus placebo after complete resection of high-risk stage III melanoma (EORTC 18071): a randomised, double-blind, phase 3 trial. Lancet Oncol 2015; 16: S22–S30.

71. Hauschild A, Dummer R, Wolchok JD, Schmidt H, et al. Adjuvant ipilimumab versus placebo after complete resection of high-risk stage III melanoma: long-term follow-up results of the European Organisation for Research and Treatment of Cancer 18071 double-blind phase 3 randomised trial. Eur J Cancer 2019; 119: 1–10.

72. Weber JS, Mandala M, Del Vecchio M, Gogas H, Arance AM, Cowey CL, et al. Adjuvant therapy with nivolumab (NIVO) versus ipilimumab (IPI) after complete resection of stage III/IV melanoma: Updated results from a phase III trial (CheckMate 238). J Clin Oncol 2018; 36: 9502.

73. Maio M, Lewis K, Demidov L, Mandala M, Bondarenko I, Ascierto PA, et al. Adjuvant vemurafenib in resected, BRAF(V600E) mutation-positive melanoma (BRIM8): a randomised, double-blind, placebo-controlled, multicentre, phase 3 trial. Lancet Oncol 2018; 19: 510–520.

74. Dahan O, Grob JJ, Wolchok JD, Gogas H, Arance AM, et al. Patterns of histologic response to neoadjuvant combination therapy with pembrolizumab and nivolumab in metastatic melanoma. Sci Transl Med 2015; 7: 279ra104.

75. Amaria RN, Prieto PA, Tetzlaff MT, Reuben A, Andrews MC, Ross MI, et al. Neoadjuvant plus adjuvant dabrafenib and trametinib versus standard of care in patients with high-risk, surgically resectable melanoma: a single-centre, open-label, randomised, phase 2 trial. Lancet Oncol 2018; 19: 181–193.

76. Long GV, Saw RPM, Lo S, Nieweg OE, Shannon KF, Gonzalez-Moreno C, et al. Neoadjuvant dabrafenib combined with trametinib for resectable, stage IIIIB-C, BRAF(V600) mutation-positive melanoma (NeoCombI): a single-arm, open-label, single-centre, phase 2 trial. Lancet Oncol 2019; 20: 961–971.

77. Slot S, Zager JS, Kudchadkar RR, Messina JL, Benedict JG, Gonzalez-Moreno C, et al. BRAF inhibition for advanced locoregional BRAF V600E mutation-positive melanoma: a potential neoadjuvant strategy. Melanoma Res 2016; 26: 83–87.

78. Zippel D, Markel G, Shapiro-Frommer R, Ben-Betzalel G, Goitzen D, Ben-Ami E, et al. Perioperative BRAF inhibitors in locally advanced stage III melanoma. J Surg Oncol 2017; 116: 856–861.

79. Ergoul Z, Khushalani NI, Rich J, Sarnaik A, Zager JS, Markowitz J, et al. Patterns of histologic response to neoadjuvant targeted therapy in patients with BRAF mutant melanoma. J Clin Oncol 2017; 35: 9584.