Immunotherapy Bridge 2017 and Melanoma Bridge 2017: meeting abstracts

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Immunotherapy bridge 2017—Keynote speaker presentations

**System biology session: immunology**

**K1**
Genomics and immunotherapy in lung cancer: tumor mutation burden, mutations affecting antigen presentation, immune recognition, and genome integrity
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*Journal of Translational Medicine* 2018, 16 (Suppl 1):K1

Immunotherapy approaches targeting the PD-1 pathway have shown some clinical benefits in a fraction of patients with lung cancer, but expression of PD-L1 has proven to be an imperfect biomarker of efficacy. Recent studies have shown that tumor mutation burden (TMB) is also correlated with outcome and that it appears to be independent of PD-L1. TMB, however, only indirectly measures the number of neoantigenic peptides presented on tumor cell surface class I MHC and predicted MHC matches may be an even better predictor of benefit. In addition, mutations in genes such as LKB1 may modulate the immune response, and mutations in the antigen presentation pathway may block it altogether. Mutations in DNA repair pathway genes may increase the number of potential neoantigens. Thus in depth analysis of tumor somatic genomics could lead to better patient selection for immunotherapy.

**SITC session—evolving topics in cancer immunotherapy**

**K2**
Discussion about evolving topics in cancer immunotherapy
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*Journal of Translational Medicine* 2018, 16 (Suppl 1):K2

Dendritic cell vaccines have been used to promote anti-tumor immune responses in over 200 clinical trials. While clinical responses have been limited, DC could play a critical role in promoting T cell infiltration into tumors which could be capitalized on with checkpoint blockade. Combination approaches including DC could also lead to improved outcomes. We tested an antigen-engineered DC vaccine with or without subsequent IFNα in melanoma. Trial results and immune biomarker data will be presented.

**Immunotherapy in oncology: data from clinical trials**

**K3**
Consequences of rapid evolution of adjuvant therapy landscape in melanoma
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*Journal of Translational Medicine* 2018, 16 (Suppl 1):K3

The spectacular outcomes of the phase III trials regarding nivolumab versus ipilimumab in fully resected stage IIIB/C-IV and of the combination of dabrafenib (D) plus trametinib (T) in BRAF mutant stage III patients demonstrate that effective treatments in advanced melanoma are also highly effective in the adjuvant setting. In 2016 an overall survival benefit with adjuvant high dose ipilimumab was demonstrated, and the EORTC1325 trial comparing pembrolizumab versus placebo will complete the picture early 2018. Toxicity profiles are in line with the experience in advanced melanoma, i.e. favorable for the anti-PD1 agents and for D+T, and problematic for ipilimumab. The 2017 outcomes are practice changing and put an end to the use of interferon and ipilimumab. In countries with only access to interferon, its use can be restricted to patients with ulcerated melanoma, based on the individual patients data meta-analysis recently published. Because of the results of the MSLT2 trial, completion lymph node dissection (CLND) will decrease sharply leading to a lack of optimal prognostic information. Prognosis in SN-positive stage IIIB/B patients is extremely heterogeneous with 5 year survival rates varying from 90% to 40%, and depends mostly on number of positive nodes identified by CLND. This information is crucial for clinical decision making. How to guarantee optimal staging information needs to be discussed urgently. Further improvements of adjuvant therapies will have to address all these questions as well as the exploration of neo-adjuvant use of active drugs and combination approaches. Important paradigm shifts in the management of high risk melanoma patients are upon us.

**Immunotherapy bridge 2017—oral presentations**

**System biology session: immunology**

**O1**
Interferon-gamma (INFγ), a predictive factor of response to immune checkpoint blockade (ICB) in melanoma and non-small cell lung cancer (NSCLC)
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The spectacular outcomes of the phase III trials regarding nivolumab versus ipilimumab in fully resected stage IIIB/C-IV and of the combination of dabrafenib (D) plus trametinib (T) in BRAF mutant stage III patients demonstrate that effective treatments in advanced melanoma are also highly effective in the adjuvant setting. In 2016 an overall survival benefit with adjuvant high dose ipilimumab was demonstrated, and the EORTC1325 trial comparing pembrolizumab versus placebo will complete the picture early 2018. Toxicity profiles are in line with the experience in advanced melanoma, i.e. favorable for the anti-PD1 agents and for D+T, and problematic for ipilimumab. The 2017 outcomes are practice changing and put an end to the use of interferon and ipilimumab. In countries with only access to interferon, its use can be restricted to patients with ulcerated melanoma, based on the individual patients data meta-analysis recently published. Because of the results of the MSLT2 trial, completion lymph node dissection (CLND) will decrease sharply leading to a lack of optimal prognostic information. Prognosis in SN-positive stage IIIB/B patients is extremely heterogeneous with 5 year survival rates varying from 90% to 40%, and depends mostly on number of positive nodes identified by CLND. This information is crucial for clinical decision making. How to guarantee optimal staging information needs to be discussed urgently. Further improvements of adjuvant therapies will have to address all these questions as well as the exploration of neo-adjuvant use of active drugs and combination approaches. Important paradigm shifts in the management of high risk melanoma patients are upon us.
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Journal of Translational Medicine 2018, 16 (Suppl 1):O1

Background: PD-L1 is up-regulated via IFNG in a STAT1- and NFkB-dependent manner. We explored whether IFNG expression in pre-treatment tumors is associated with the activity of ICB in NSCLC and melanoma patients. The role of inflammation-associated transcription factors STAT3, IKBKE and STAT1 was also examined.

Methods: Total RNA from 17 NSCLC and 21 melanoma patients, was analyzed by qRT-PCR. STAT3 and Rantes, YAP1 and CXCL5, DNN1T1, RIG1 and TET1, EOMES, INFg (encoding for IFNG), PD-L1 and CTLA4, IKBKE and NFATC1 mRNA were examined. PD-L1 protein expression in tumor and immune cells and stromal infiltration of CD8+ T cells were also evaluated.

Results: 17 previously treated NSCLC patients received nivolumab; 71% lung adenocarcinoma, 71% male, 53% smokers, 35% KRAS mutant, 88% EGFR wild-type (wt). 21 previously treated melanoma patients received pembrolizumab; 67% male, 67% BRAF wt. PFS to nivolumab was significantly longer in NSCLC patients with high vs low IFNG expression (5.12 vs 2 months, p = 0.0124). PFS to pembrolizumab was significantly longer in melanoma patients with high vs low IFNG expression (4.99 vs 1.86 months, p = 0.0099). Significantly longer OS was observed for melanoma patients with high vs low IFNG expression (not reached vs 3.10 months, p = 0.0183). There was a trend for longer OS for NSCLC patients with high vs low IFNG expression (10.15 vs 4.86 months, p = 0.0687). A survival plateau was only observed for patients with high IFNG levels: OS31m, 60% vs 0% for melanoma, and 15% versus 0% for NSCLC. Clinical benefit (CB)(objective response or stable disease) was observed in 71.4% of NSCLC patients with high IFNG levels versus 20% for patients with low IFNG. The tumor and immune cells did not affect the outcome to ICB. IKBKE was positively correlated with IFNG and PD-L1 expression (NSCLC Spearman’s ρ = 0.58 and 0.65; melanoma Spearman’s ρ = 0.61 and 0.59), and STAT3 expression was loosely anticorrelated with PD-L1 expression (NSCLC Spearman’s ρ = −0.21; melanoma Pearson’s ρ = −0.01). The rest of the biomarkers explored did not affect the outcome to immunotherapy.

Conclusions: NSCLC and melanoma patients with intermediate/high IFNG mRNA expression exhibited longer PFS and OS and higher disease control rates with anti PD-1 therapies even when the levels of PD-L1 expression were low.

O2 Biomarkers for cancer immunotherapy: predicting the immune resistance through gene expression profile
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Journal of Translational Medicine 2018, 16 (Suppl 1):O2

Background: Surgery, radiation and chemotherapy have been long considered the three pillars of cancer care. A fourth pillar, immunotherapy, recently expanded providing an exciting new treatment option for many patients. In this regard, monoclonal antibodies blocking PD-1/PD-L1 immune checkpoint have shown promising clinical results [1].

Despite these very encouraging results, the majority of patients do not respond to immunotherapy regimens as monotherapy, leading to an urgency to identify biomarkers that accurately predict which patients will benefit from treatment and potentially actionable mechanisms of resistance in order to set successful combinatorial approaches [2]. To this end, gene-expression profiling has been here applied on lesions derived from patients with renal cell carcinoma (RCC), melanoma and classical Hodgkin lymphoma (CHL) thus unveiling novel and paradoxical relations between cancer and immune system leading to immunotherapy resistance.

Method: RNA was isolated from (i): 13 formalin-fixed, paraffin-embedded (FFPE) pre-PD-1 treatment tumor biopsies derived from RCC patients; (ii): 10 FFPEs regressing/progressing cutaneous metastases derived from on autopsy case of melanoma; (iii): 24 FFPEs derived from CHL Epstein Barr virus (EBV) positive (+) and negative (−) patients. RNA was subjected to whole genome microarray and multiplex quantitative (q) RT-PCR.

Results: In renal cell carcinoma, gene expression profile highlighted metabolic and immunologic molecules to be associated with the effective response to immunotherapy with anti- PD-1 blockade [3]. In melanoma, transcriptional signatures mostly associated with epithelial to mesenchymal transition (EMT) and accumulation of neutrophils were found to be associated with PD-1 blockade therapy resistance [4]. In CHLs, results revealed a dichotomous cellular and cytokine immune milieu in EBV+ vs EBV− CHL [5]. Particularly, EBV+ tumors displayed a T helper 1 (Th1) profile while EBV-tumors manifested a pathogenic Th17 profile and ongoing engagement of the interleukin-23 (IL-23)/IL-17 axis [6]. These findings suggest that drugs blocking the IL-23/IL-17 axis, may enhance the therapeutic impact of immunotherapy in EBV−CHL.

Conclusions: Many pathways might determine the clinical response to immunotherapies in cancer patients, thus suggesting that in the evaluation of biomarkers associated with response to immunotherapy, all intersections between immunological, genetic and tissue specific factors must be evaluated. Merging together the usage of high-throughput screenings, bioinformatic analysis and immune biology assays might be necessary to establish a framework for describing the diversity of these interactions with the aim to focus on features that help guide immunotherapeutic treatment choices on an individual basis (i.e. personalized medicine).

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O3 4SC-202 induces inflamed tumor microenvironment, strongly enhances tumor infiltration with cytotoxic T cells and primes tumor microenvironment for anti-PD-1/PD-L1 therapy
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Journal of Translational Medicine 2018, 16 (Suppl 1):O3
**Background:** Various HDAC inhibitors were described as beneficially affecting anti-tumoral immune response. Although different HDAC inhibitors were investigated in syngeneic tumor models, the mode of anti-tumoral action is not yet fully understood. Here, we analyzed the anti-tumoral mode of action (MoA) of 4SC-202, an orally available clinical stage epigenetic small molecule inhibitor targeting histone deacetylases (HDAC) class I. To ensure that the conclusions would be relevant for the clinical situation we used a clinically equivalent dose-regime.

**Materials and methods:** Anti-tumoral efficacy and the impact on tumor microenvironment (TME) were analyzed in syngeneic colorectal CT26 and C38 models in immunocompetent BALB/c or in nude/irradiated BALB/c mice. A broad spectrum HDAC inhibitor was used for comparison. Transcriptome analysis was performed by RNA-Seq, and the composition of immune cell subpopulations was determined by flow cytometry.

**Results:** 4SC-202 treatment significantly inhibited growth of CT26 and C38 tumors. A competent immune system was apparently necessary for the anti-tumoral effect of 4SC-202 since its tumor-reducing effect was lost in immunocompromised mice. 4SC-202 treatment increased IFN-γ and chemokine expression, and reduced pro-inflammatory IL-1 and IL-23 in the TME of CT26 tumors. Detailed analysis revealed that 4SC-202 increased the number of cytotoxic CD8+ T cells (CTLs) in TME of both, T cell-inflamed C38 as well as of non-T cell-inflamed CT26 tumors without affecting the number of CTLs in blood. In contrast, a broad-spectrum HDAC inhibitor tested in the same model demonstrated anti-tumoral efficacy but did not affect the number of CTLs in tumors demonstrating that HDAC inhibitors employ different MoAs for their anti-tumoral response and that the effect on CTLs is not attributed to HDAC inhibition in general. Since the T cell abundance is pre-requisite for the efficacy of PD1/PD-L1 blockade, combinations of 4SC-202 with anti-PD-1/anti-PD-L1 antibodies were tested in C38 and CT26 models, respectively. The combined treatment was more efficacious than monotherapies and resulted in significantly longer survival in both models with 55% tumor-free animals in C38 model.

**Conclusions:** 4SC-202 already demonstrated a favorable safety profile in a phase I clinical trial with relapsed or refractory hematological malignancies with two objective responses (1 CR, 1 PR) and disease stabilizations in several patients. 4SC-202’s immune priming capacity offers further options for clinical development of 4SC-202 in combination with various cancer immunotherapy approaches. Combination of 4SC-202 with PD-1 blockade will be evaluated in a Phase Ib/II clinical study in advanced cutaneous melanoma patients refractory/non-responding to treatment with anti-PD-1 antibodies (‘SENSITIZE’, NCT03278665).

**Immunotherapy beyond melanoma**

**O4**

**A new triple negative breast cancer (TNBC) murine model for in vivo preclinical immunotherapies**

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**Journal of Translational Medicine 2018, 16 (Suppl 1):O4**

**Background:** Triple Negative Breast Cancers (TNBCs), lacking hormone receptors and HER2, are highly metastatic and chemoresistant and metastatic events (mBC) are the most common cause of death in women. Tumour microenvironment (TME) is a complex network of cells that supports tumorigenesis and metastatic spread. Among the immune cells of TME, Tumour-Associated-Macrophages (TAMs) are the most abundant in BC by regulating invasion, metastases and chemoresistance. TAMs can acquire distinct phenotypes in response to different signals. M2-polarized-TAMs have immunosuppressive activities by expressing inflammatory molecules. Prune-1 belongs to DHX (Asp-His-His) phosphoesterase superfamily with an exopolyphosphatase activity [2, 8]. The overexpression of Prune-1 is correlated with metastases and poor prognosis in several tumours including BC [8]. Prune-1 has also been found to induce Epithelial-Mesenchymal-Transition (EMT) and metastatic dissemination through the enhancement of canonical TGF-β signalling by counter-balancing its inhibition operated by NM23-H1 [3, 4]. Furthermore, we also have evidences that lung cancer progression is driven by Prune-1 via canonical WNT signalling in autocrine and paracrine manner via Wnt3a secretion [1].

**Results:** We identified Prune-1 with the ability to recruit and polarize TAMs toward a pro-tumorigenic M2-phenotype within the TME of TNBC using a double Genetically Engineered Mouse (GEM) model of TNBC over-expressing both Prune-1 and Wnt-1 in mammary glands (MMTV-Prune-1/Wnt1); generated through the use of vectors construct containing the human transgene cDNAs under the control of Mouse Mammary Tumour Virus (MMTV) promoter. These novel Genetically Engineered Mouse (GEMs) model of TNBC (MMTV-Prune-1/WNT1) develop BCs with 100% penetrance between months 2–3 of life (starting at birth), fully developed and they always generate lung metastases, while the single transgenic MMTV-Wnt1 TNBC models [5, 7] is not able to make them. These GEMs were then crossed with the receptor 2 of VEGF (VEGFR2) promoter driving firefly luciferase gene expression. Results are indicating that once these recombinant animals MMTV-Wnt1 or MMTV-Prune-1/Wnt1 develop tumours activating VEGF then these initiating tumorigenic cells can be visible by in vivo bioluminescence imaging (BLI) luciferase technology. Studies in primary cells derived from the BC generated by these GEM models, indicate that the over-expression of Prune-1 is responsible for the activation of intracellular pathways (i.e. TGF-β, FAK and NF-κB) and for both the activation and polarization of macrophages in vitro shown by the activation of JAK-STAT3 and NF-κB signalling cascades and the increase of inflammatory cytokines (e.g. Arg1, iNOS, MPP9 and IL1β) in those macrophages treated with conditioned media derived from MMTV-Prune-1/Wnt1 primary cells. This thus confirms Prune-1 able to polarize TAMs toward an M2-phenotype.

**Conclusions:** We generate a TNBC murine model with lung metastases which can be monitored by in vivo imaging (BLI) technology. This GEM model can be an useful source for immunotherapy trials being a model of enhancement of M2-TAMs polarized cells within the TME in primary tumour and lung metastases. These results are of impact for immunotherapy for studies with new check-points inhibitors with activities against these specialized cells.

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Characterization of melanoma patients with brain metastases diagnosed between 2014–2016, in one center

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O5 Characterization of melanoma patients with brain metastases diagnosed between 2014–2016, in one center

Background: The treatment of patients suffering from melanoma brain metastasis is challenging. Combination of local and systemic therapies is under evaluation in several ongoing clinical trials. Moreover, treatment sequence needs further optimization and therefore standards for the management of brain metastases in melanoma patients do not exist so far.

Methods: After approval of the ethical commission, we conducted a retrospective study including 168 patients diagnosed with melanoma brain metastases between 2014 and 2016 and treated with local and/or systemic therapies. The cut-off date for data collection was April 2017. Overall survival was analyzed using a Kaplan–Meier estimator. Results: The median follow-up since the first melanoma diagnosis was 61.8 months [23.0–80.75] and 8.59 months [3.0–12] since the first diagnosis of brain metastasis. As of the cut-off date, 39% of the patients were still alive. The median melanoma specific survival, defined as the time between melanoma diagnosis and last observation or death, was 63 months [25.0–129]. The median overall survival (OS) for the all population, defined as the time between brain metastasis diagnosis and last observation or death, was 9 months [4.0–22.0]. For patients treated with immunotherapy as first systemic therapy, the median OS was 13 months (95% CI 7.65–18.35 months) and 11 months (95% CI 6.55–15.46 months; p = 0.005) for those treated with targeted therapy. When the type of first line local therapy is analyzed, the median OS was 22 months (95% CI 11.24–31.76 months) for patients treated with surgery or stereotactic radiotherapy and 6 months (95% CI 4.36–7.64 months; p = 0.0001) for patients treated with whole brain radiation. The best results were obtained when both systemic and local therapies were combined in a 4 weeks interval, but this was not significant (p = 0.061). In patients with BRAF mutation, longer median OS was observed in patients treated with immunotherapy as first systemic therapy when compared to targeted therapy. The median OS was not reached in the first group and was 11 months (95% CI 6.54–15.45 months; p = 0.004) in the second group.

Conclusion: The availability of new therapies increased OS of patients with brain metastases, in comparison with historical controls (9 months vs 5 months). Immunotherapy as first systemic therapy was associated with the best outcomes, including in patients harboring BRAF mutation.

O6 A novel translational research tool for the development of predictive signatures of immunotherpay response

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Journal of Translational Medicine 2018, 16 (Suppl 1):O6

The Tumor Inflammation Signature (TIS) is an 18 gene biomarker of a suppressed adaptive immune response within tumor which measures four key areas of biology—antigen presentation, T/NK cell abundance, INFγ signaling, and T cell exhaustion. The TIS is currently under evaluation in three clinical trials to predict immune response to pembrolizumab, and may have broad utility to predict response to other immune checkpoint inhibitors. The TIS has been embedded into the NanoString® IO 360 panel—a 770 gene expression panel allows for the parallel assessment of additional mechanisms of immune-evasion in the RUO setting using a single 5 μm FFPE tissue section. The panel contains content to characterize evasion in the context of an inflamed tumor phenotype (such as additional checkpoints inhibitors or suppressive immune cell populations) as well as in the context of an “immune excluded” or “immune desert” tumor microenvironment phenotype (such as activation of oncogenic pathway affecting immune cell trafficking or intrinsic alteration of the antigen presentation process). The IO 360 panel enables the development of diagnostic tests that will select populations that respond to novel and existing immunotherapies as well as combination therapies based on the parallel assessment and integration of multiple mechanisms of immune evasion in a single assay.

SITC session—evolving topics in cancer immunotherapy

O7 Mechanisms of primary resistance to cancer immunotherapies

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Background: Although much clinical progress has been made in harnessing the immune system to recognize and target cancer, there
is still a significant lack of an understanding of how tumors evade immune recognition and the mechanisms that drive tumor resistance to both T cell and checkpoint blockade immunotherapy. Our objective is to understand how tumor-mediated signaling through inhibitory receptors, including PD-1, combine to affect the process of T cell recognition of tumor antigen and activation signaling, with the goal of understanding the basis of resistance to PD-1 blockade and the potential identification of new molecular targets to enable T cells to overcome dysfunction mediated by multiple inhibitory receptors.

Methods and results: We show that Lck activity affects T cell sensitivity and influences the probability of inducing effector function [1]. Under non-activating conditions, Lck and Shp-1 phosphorylation and activity vary based on CD8+ memory T cell phenotype. Shp-1 interaction with Lck under non-activation conditions can also vary, as suggested by our results showing decreased Shp-1 S591 phosphorylation, which affects Shp-1 localization and correlates with increased Shp-1 colocalization with Lck. Further, we showed that Shp-1 directly influences Lck activity under non-activating conditions, as inhibition of Shp-1 leads to increased Lck activity. Importantly, inhibition of Shp-1, a major mediator of PD-1 signaling, targeting CD28 and Lck [2], prior to activation leads to increased T cell cytotoxic effector function. Our proteomics-based analysis of patient T cells identified both mediators of PD-1 signaling and signaling components and pathways associated with blockade resistance. It has generally been thought that TCR and CD8 binding depend mainly on their ectodomain interactions with pMHC. We have shown, however, that Lck-CD8 binding [3] and Lck activity [4] are required for upregulated CD8 binding to pre-bound TCR-pMHC complex. Therefore, the cytoplasmic associations of Lck with CD8 and Zap-70, as well as CD3 with Zap-70 may influence formation and stability of the TCR-pMHC–CD8 complex. To determine the mechanistic basis of PD-1 inhibition of TCR–pMHC–CD8 binding, we utilized 2D affinity combined with Biomembrane Force Probe (BFP) measurements [5, 6] and showed that PD-1 directly suppresses TCR–pMHC–CD8 binding. Our data also revealed that TCR-pMHC binding was independent of PD-1-PD-L1, but TCR–pMHC–CD8 binding was suppressed by PD-1-PD-L1 binding demonstrating negative cooperativity, as fewer bonds formed than the sum of bonds formed by each interaction alone.

Conclusions: Together, our results show that the activities of TCR-proximal signaling components affect T cell mechanosensing and sensitivity at the earliest stages of antigen recognition and are influenced by PD-1. Targeting these interactions may enhance tumor-specific T cell sensitivity for cancer immunotherapy and understanding the basis of resistance to PD-1 blockade to potentially allow identification of new molecular targets to enable T cells to overcome dysfunction mediated by multiple inhibitory receptors.

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09 Role of TGFβ superfamily members in hindering the pro-inflammatory effects of radiotherapy
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Background: Transforming Growth Factor-beta (TGFβ) and activin A (actA) are TGFβ superfamily members with overlapping functions in many processes including regulation of inflammation and immunity. We have recently shown that in situ vaccination by local tumor irra-
diation is hindered by activation of latent TGFβ [1]. Intriguingly, TGFβ blockade enhanced activation of dendritic cells and T-cell priming, but it increased (rather than reduced) intratumoral regulatory T cells (Tregs). We have recently found that actA release by breast cancer cells is enhanced by radiotherapy (RT). Interestingly, prolonged exposure to TGFβ inhibitors also resulted in actA upregulation, consistent with a previously described compensatory mechanism. Here we hypothe-
sized that actA and TGFβ regulate RT-induced anti-tumor immunity.

Methods: Secretion of actA by untreated and irradiated 4T1 mouse carcinoma cells was quantified by ELISA. Transwell co-culture was used to assess the ability of cancer cell-derived actA to promote the conversion of naive CD4+ T cells into Tregs. 4T1 cell derivatives engineered to express a tetracycline-inducible shRNA specific for actA (4T1ΔactA) or non-silencing (4T1ΔshNS) were generated and injected s.c. to syngeneic BALB/c mice (day 0). ActA knockdown was induced by systemic doxycycline administration at day 8. TGFβ-neutralizing 1D11 or isotype control antibodies were given i.p. every other day starting on day 12. RT was delivered to the primary tumor in 6 Gy fractions on 5 consecutive days beginning on day 13. Mice were followed for tumor growth or euthanized on day 22 for analysis.

Results: TGFβ blockade improved RT-mediated tumor control, an effect mediated by T cells. However, tumor recurred. Notably, ActA KD or 1D11 increased intratumoral Tregs (Control: 11%; 1D11: 26%; shActA: 21%) and enhanced Tregs infiltration induced by RT (RT: 15%; RT+1D11: 27%; RT+shActA: 50%). When both TGFβ and actA were blocked Tregs significantly decreased in both untreated (1D11+shActA: 13%) and RT-treated tumors (RT+1D11+shActA: 8% of Tregs). Tumor-specific IFNγ production by CD8+ T cells was sig-
nificantly higher in RT+1D11+shActA-treated mice compared to RT+1D11 (*p) and RT+shActA (**p). ActA KD in mice treated with RT+1D11 reduced tumor recurrence and improved survival (RT+1D11 vs RT+1D11+shActA **p; RT+shActA vs RT+1D11+shActA ***p).

Conclusion: Data indicate that both TGFβ and actA impair RT-induced anti-tumor immune responses. Concomitant inhibition of actA and TGFβ is required for optimal in situ vaccination by RT.

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1. Vanpouille-Box C, et al. TGFβ is a master regulator of radiation therapy-induced anti-tumor immunity. Cancer Res 2015;75(11):2322–42.
Background: Despite the emergence of active new systemic therapies, metastatic melanoma remains a clinically challenging form of skin cancer. The renin-angiotensin system (RAS) is a major physiological regulatory pathway mediated by angiotensin II (AngII) via two receptor subtypes, AT1R and AT2R. The role of the AT1R in melanoma is unexplored in melanoma.

Materials and methods: We investigated the involvement of the two principal angiotensin receptors in a panel of melanoma cell lines, grown as described previously. Primary cultures of brain metastatic melanomas were established from fresh tumour surgical tissues. The selective AT1R inhibitor Losartan and the highly selective AT2R agonist Y6All were developed as described. Demethylation experiments using azacytidine and trichostatin were done as described. TaqMan® probes were used for gene expression analysis. Transfectants with AT1R ORF were analysed for knockdown of AT1R by qPCR and WB. Cell proliferation and clonogenic assays were assessed by standard techniques. The role of AT1R in tumour angiogenesis was investigated in hCMEC/D3 grown in CM collected from PMWK cells treated with AngII alone or in combination with Losartan and PD123319.

Results: Antagonism of AT1R using the Losartan or shRNA-mediated knock-down in melanoma cell lines expressing AT1R resulted in acquisition of the ability to proliferate in serum-free conditions, implying that AT1R has a negative growth-regulatory function in melanoma. Consistent with this, ectopic expression of AT1R in cell lines lacking endogenous expression inhibits proliferation irrespective of the presence of AngII implying a ligand-independent suppressor function for AT1R. Treatment of melanoma cell lines expressing endogenous AT1R with either AngII or the AT2R-specific agonist Y6All induces proliferation in serum-free conditions. Conversely, the AT2R-specific antagonists PD123319 and EMA401 inhibit melanoma growth and angiogenesis and potentiate inhibitors of BRAF and MEK.

Conclusions: Our results indicate that the RAS has both oncogenic and tumour suppressor functions in melanoma. Pharmacological inhibition of AT1R may have therapeutic effects in melanomas expressing this receptor and AT1R methylation in serum may serve as a biomarker of metastatic melanoma.

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O10 Functional genomics to identify germline markers of melanoma immunotherapy efficacy and toxicity
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Background: Approximately 40% of metastatic cutaneous melanoma (CM) patients do not respond to the current immunotherapy (IT) regimen, pointing to other, yet unknown factors conferring IT resistance. In addition, > 60% of patients from single-line or combined treatment (COMBO) regimens present severe immune related adverse events (irAEs). In this study we have developed a novel genomic approach interrogating expression quantitative trait loci (eQTLs) to explore weather germline genetic variation can serve as novel personalized determinant of immunotherapy response and toxicity.

Methods: By interrogating the genome wide expression data and SNP array datasets of healthy twin cohort (MuTHER), we have identified 85 eQTLs most significantly associated with the expression of 265 immune genes. Using the MassARRAY system, the 85 SNPs were genotyped in 138 anti-CTLA-4 treated patients, 87 PD-1 treated patients, and 69 patients from combined (COMBO) treatments, collected from multi-institutional collaborations. To test the association of SNPs with IT response and irAEs, logistic regression analyses were performed for each treatment group adjusting by demographic and clinical covariates.

Results: We found significant associations with COMBO IT resistance for and eQTL in IL10/IL19 (OR = 4.249, p = 0.0167), which we have recently identified for association with melanoma survival and which, interestingly, is an established locus associated with the risk of several autoimmune diseases. Additionally, we also identified eQTLs that are associated with IT sensitivity; IL1-β with resistance to anti-CTLA-4 and SPI1 with resistance to anti-PD-1. Interestingly, genomic scan of 85 eQTLs has identified novel loci predictive of severe autoimmunity and site specific irAEs in patients treated with COMBO or single-line anti-CTLA4 IT.

Conclusions: In this study, we report that eQTLs from IL19/IL10 locus, previously shown to predict autoimmunity risk and CM survival, is also a surrogate marker of response to COMBO IT, indicating a strong relationship between interleukin pathways and tumor immunogenicity. Novel loci have been found as predictive markers for autoimmune toxicity, in patients treated with COMBO and anti-CTLA4 IT. This is a first evidence that immunomodulatory pathways modulated by germline genetic variation can impact susceptibility to irAEs as well as IT efficacy. Currently, a large scan is underway using genome-wide genetic screens to further test the functional validity of these findings in a large collaborative setting.

Focus on health cost assessments and pharmacovigilance in immunotherapy

O11 Comparison between new therapies of metastatic and/or unresectable melanoma with BRAF V600E/K mutations: cost–benefit assessment
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Journal of Translational Medicine 2018, 16 (Suppl 1):O10

Background: Approximately 40% of metastatic cutaneous melanoma (CM) patients do not respond to the current immunotherapy (IT) regimen, pointing to other, yet unknown factors conferring IT resistance. In addition, > 60% of patients from single-line or combined treatment (COMBO) regimens present severe immune related adverse events (irAEs). In this study we have developed a novel genomic approach interrogating expression quantitative trait loci (eQTLs) to explore whether germline genetic variation can serve as novel personalized determinant of immunotherapy response and toxicity.

Methods: By interrogating the genome wide expression data and SNP array datasets of healthy twin cohort (MuTHER), we have identified 85 eQTLs most significantly associated with the expression of 265 immune genes. Using the MassARRAY system, the 85 SNPs were genotyped in 138 anti-CTLA-4 treated patients, 87 PD-1 treated patients, and 69 patients from combined (COMBO) treatments, collected from multi-institutional collaborations. To test the association of SNPs with IT response and irAEs, logistic regression analyses were performed for each treatment group adjusting by demographic and clinical covariates.

Results: We found significant associations with COMBO IT resistance for and eQTL in IL10/IL19 (OR = 4.249, p = 0.0167), which we have recently identified for association with melanoma survival and which, interestingly, is an established locus associated with the risk of several autoimmune diseases. Additionally, we also identified eQTLs that are associated with IT sensitivity; IL1-β with resistance to anti-CTLA-4 and SPI1 with resistance to anti-PD-1. Interestingly, genomic scan of 85 eQTLs has identified novel loci predictive of severe autoimmunity and site specific irAEs in patients treated with COMBO or single-line anti-CTLA4 IT.

Conclusions: In this study, we report that eQTLs from IL19/IL10 locus, previously shown to predict autoimmunity risk and CM survival, is also a surrogate marker of response to COMBO IT, indicating a strong relationship between interleukin pathways and tumor immunogenicity. Novel loci have been found as predictive markers for autoimmune toxicity, in patients treated with COMBO and anti-CTLA4 IT. This is a first evidence that immunomodulatory pathways modulated by germline genetic variation can impact susceptibility to irAEs as well as IT efficacy. Currently, a large scan is underway using genome-wide genetic screens to further test the functional validity of these findings in a large collaborative setting.

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Background: Incidence of melanoma continues to rise, and the mortality associated with unresectable or metastatic melanoma remains high. B-RAF targeted therapy has been established as a treatment standard for patients who have metastatic melanoma with activating BRAF mutations and recently a regimen combining a B-RAF inhibitor with a MEK inhibitor has been associated with a higher response rate and longer duration of response, as compared with anti-BRAF monotherapies. These drugs are subject monitoring by Italian Management Entry Agreements (MEAs). The objective of our study is to evaluate the cost–benefit evaluation of anti-BRAF and anti-MEK agents alone or in combination to calculate the average of pharmacological cycles performed, the number of progressions and the percentage of partial and/or complete remission percentage Treatments That Meet MEAs.
Methods: The clinical, therapeutic and diagnostic data of each patient were obtained from the hospital databases and the monitoring registry by Italian Medicines Agency, that contains data for monitoring patients who are receiving this therapies. The report of this register includes the medicines that are registered and some of the outcomes that were being monitored, for example, the number of treated patients, the patients that have finalized the treatment and the reasons for stopping the treatment. In these monitoring, we are included all patients with advanced melanoma with B-RAF V600E/K mutation treated between 2012 and 2016. Result: 112 patients with non-resectable metastatic melanoma were treated from 2012 to 2016, of which 68 were treated with monotherapy (65 with Vemurafenib and 3 with Dabrafenib), 44 with BRAF and MEK-inhibitor associations (12 Vemu/Cob and 32 Dab/Tram). Patients performed on average 12 cycles of Vemurafenib, 3 cycles of Dabrafenib, 5 cycles of Vemu/Cob and 6 cycles of Dab/Tram. Compared to the MEAs, over half of the therapies, on average 60%, monotherapy or association, meet the expectations described.

Conclusion: Higher PIS therapies are monotherapy with Vemurafenib (4,4) and the Dabr/Tram (3,26) association, which on average patients take the drug for more time following the Vemurafenib monotherapy (12 cycles), exceeding the MEAs expectations.

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O13 The drug day an effective strategy for the containment of immunotherapy costs
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Journal of Translational Medicine 2018, 16 (Suppl 1):O12

Background: The Centralized Unit for Handling Antineoplastic of National Cancer Institute “G. Pascale” of Naples planned a strategy for the use of high cost cancer drug Opdivo®. In order to reduce the therapy cost we decide to dedicate for Opdivo® a drug day to optimize the use of drug vials with reduction of waste and/or optimal use of the residues. The aim of this work was to report the analysis of drug consumption and cost from March 2016 when it was officially authorized as a hospital drug and it was introduced into clinical practice to August 2017.

Materials and methods: The drug day was organized in accord with Opdivo® prescribed physicians. The number of patients treated, the individual patient’s treatment line, consumption and cost data relating to the period under review, were collected from the hospital database. In primis, it was performed an assessment of the total consumption and cost per year and then a comparison exercise was made between the number of drug vials really used and those that should be used without the drug day and, the comparison between the real cost incurred and the hypothetical cost out of drug day.

Results: From March 2016 to August 2017 the pharmacy staff set 1872 preparation for 211 patients of Opdivo®. The drug was administered for 68 patients as first-line treatment, for 116 patients as second-line treatment and finally for 27 patients as third-line treatment. Given the considerable number of patients and considering that Opdivo® stability is 24 h, 2 consecutive days per week were dedicated for treatment as drug days with a range of 15–20 preparations per day. Comparing the used drug in drug day to a hypothetical daily preparation we saved 36,860 mg (approximately 367 vials of 100 mg) with an economy of €435,943,22. Noteworthy, the overfill of injecting drug vials, corresponding to about 10–12 mg over the declared amount of drug, as prescribed by F.U. XII Edition (2.9.17) and by FDA guidelines on the filling volume in excess of vials, allowed us to rescue 13.167 mg with an economy of €155,887,67.

Conclusions: Programming cancer therapies in a drug day was very complex and involved close cooperation between prescribed physicians, pharmacists and patients. However this strategy allows to reduce at minimum drug waste residues and, furthermore, to use overfill of the samples, which should become waste. In conclusion the drug day resulted a very effective tool for the containment of pharmaceutical costs.

O13 Adverse effects related to PD-1 inhibitor immunotherapy: an overview
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Journal of Translational Medicine 2018, 16 (Suppl 1):O13

Background: Immunotherapies, like nivolumab (Opdivo®) and pembrolizumab (Keytruda®), have changed the cancer treatment landscape. They are inhibitors of PD-1 protein, which cancer cells use to...
elude the immune system. Post-marketing surveillance of these drugs have revealed severe adverse drug reactions (ADRs). The occurrence of ADRs has high morbidity and mortality, accounting for the fifth leading cause of death in industrialised countries. EudraVigilance.org is European Medicine Agency’s (EMA) web-based Monitor System for reporting and evaluating suspected ADRs.

**Aim:** A retrospective observational study was done. The aim of this study is to investigate the ADRs occurred in patients treated with PD-1 inhibitors.

**Methods:** All ADRs associated with Opdivo® and Keytruda® reported in EudraVigilance up to 24 September 2017 were analysed for overall ADRs category for both Opdivo® and Keytruda®. A total of 10600 ADRs related to the PD-1 Inhibitors therapy were reviewed and evaluated. The proportion of ADRs (proportional reporting ratio [PRRs]) was developed for signal generation from large databases of spontaneous ADRs reports to find a statistical link between each PD-1 inhibitor and ADRs.

**Results:** A total of 10600 ADRs related to the PD-1 Inhibitors therapy have been reported and analysed in this study. Opdivo® (65.8%), Keytruda® (31.5%) and nivolumab (2.63%). 55.25% occurred in males (male/female ratio = 1.73). Only in 12.8% of ADRs reported, sex is not specified. Most ADRs occurred in patients with age between 18 and 65 years (41.6%). The higher incidence of ADRs occurred in patients of non-European Economic Area than European Economic Area, respectively 73.7% vs 26.3%.

The "Neoplasm benign malignant and unspecified" is the most reported ADRs category for both Opdivo® and Keytruda® (18.9% vs 27.7%) with a PRR = 0.67. ADRs occurred in males show PRR = 0.87. Serious but rare ADRs reported in database is Uveitis, that shows PRR = 0.61. The Uveitis incidence rate, reported in database, is 0.70 vs 0.33, respectively for Keytruda®-treated patients than Opdivo®-treated patients.

**Conclusions:** Analysis of Pharmacovigilance database is important to understand the safety of drugs in post-marketing and in real clinical practice. Patient safety is priority in care management. For this reason, oncologist, pharmacists and multidisciplinary team have to be careful about adverse reactions that could be occurred in patients treated. Limitations of this study was cases were based on spontaneous reporting which clearly suffered from underreporting. Clinical data were not available.

**Immunotherapy bridge 2017—poster presentations**

**P1**

**Excellent response to anti-PD-1 therapy in a patient with metastatic malignant melanoma: case report**

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**Journal of Translational Medicine 2018, 16 (Suppl 1):P1**

**Background:** Malignant melanoma is an aggressive cancer associated with high mortality worldwide. Many patients have locally advanced or metastatic disease at diagnosis in which case treatments were limited until a few years ago. Immune checkpoint blockade of programmed death receptor-1 (PD-1) pathway represents a recent valid treatment strategy changing the prognosis in this setting.

**Case presentation:** Here we describe the case of a 83-year-old man with many comorbidities (chronic renal failure, diabetes, hypertension) and with colorectal cancer anamnesis (total laryngectomy in 2004) and laryngeal neoplasia (total laryngectomy in 1981) who presented initially neofomation of the parotid region and a skin nodule of the temporal region were disappeared as well as neofomation of the parotid region. On October at CT imaging, the alterations to the parotid region were completely disappeared and lung metastases appeared hyperdense and cavitated. Last total body PET/TAC (on February 2017) described a single pulmonary localization to the left basal pyramid. The patient is continuing nivolumab therapy with good tolerability, with the exception of a skin rash G1-2 and he is still being in response at a distance of more than 1 year.

**Conclusions:** This case report confirms the efficacy of nivolumab in metastatic melanoma and the safety in elderly patients with polypathology also.

**Consent for publication:** The authors declare that written informed consent was obtained from the patients for publication.

**P2**

**Pharmacovigilance in melanoma immunotherapy**

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**Journal of Translational Medicine 2018, 16 (Suppl 1):P2**

**Background:** Ipilimumab, Nivolumab and Pembrolizumab recently approved by the Italian Medicines Agency for first-line therapy of melanoma. They are active on the immune system and have a toxicity related to their mechanism of action. Skin, gastrointestinal tract, liver, endocrine system are mainly affected. Additional monitoring is planned for Nivolumab and Pembrolizumab to allow rapid acquisition of new data about their safety [1, 2]. The aim of Authors is to assess whether this post-authorization monitoring has so far provided about updating the risk profile of these inhibitors of immune checkpoints.

**Materials and methods:** Suspected adverse reactions (ADRs), associated with the use of Ipilimumab, Nivolumab and Pembrolizumab in the treatment of melanoma and recorded in the Pharmacovigilance National Network in the 2015–2016 biennium, were examined and compared with the side effects reported in the Product Characteristics Summary of these drugs and the literature found in Pubmed-Medline and Cochrane Library.

**Results:** From 1 January 2015 to 31 October 2016 111 ADRs were found: 93 relate Ipilimumab, 6 Nivolumab and 12 Pembrolizumab. They were evaluated 57 reports and others could not be classified; diarrhea was the main adverse event reported and 23 ADRs were serious. Three cases only were treated with systemic immunosuppressive drugs. Analysis of adverse reactions showed that they are common side effects of three inhibitors of immune checkpoints.

**Conclusions:** The pharmacovigilance of Ipilimumab, Nivolumab and Pembrolizumab has produced an appreciable number of reports which however represent known side effects of recent melanoma immunotherapy that can be properly treated with systemic immunosuppressive drugs if recognized early [3]. Severity of adverse events has generally required hospitalization: recognition and treatment of these reactions may have been inappropriate or later. Authors believe not all health workers were joined by a clear and complete information about adverse events of these drugs, their recognition and treatment. In cooperation with the oncologist, hospital chemist is able to promote informational programs that can guarantee the knowledge necessary for an adequate approach to toxicity associated to use of these drugs. The aim is ensuring an effective and efficient health service; in this case a correct information can produce a higher yield by reducing costs associated to inappropriate therapies and excessive hospitalization.

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Bullous pemphigoid during Nivolumab therapy: a case report

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Background: Anti-programmed cell death protein-1 (PD-1)/ligand-1 (PD-L1) antibodies can induce an immune-related bullous pemphigoid (BP) [1, 2]. About its pathogenesis, it has been hypothesized that the blockade of the PD-1/PD-L1 pathway may increase autoantibody production against the hemidesmosomal protein BP180 [1]. Here we report a case of BP during treatment with nivolumab.

Case report: A 68-year-old man was admitted to our Oncology Unit in March 2015 with the diagnosis of BRAF wild-type melanoma. A CT scan performed on May 2016 showed lung metastases. Patient's medical history included hypertension. Long-standing medications included omeprazole and zofenopril. He had no relevant history of skin or autoimmune disorders and no new medications. On June 2016, nivolumab was started. On June 2017, the patient began to develop erosions and vesicles on the buccal mucosa, especially with the involvement of the lower lip. The severity of the patient's bullous dermatitis ranged from grade 1–2. A biopsy of these lesions showed eosinophilic spongiosis and a mixed dermal inflammatory infiltrate with eosinophils. Direct immunofluorescence showed linear deposition of IgG and C3 at the basement of the dermoepidermal junction, establishing a diagnosis of BP. A treatment with oral prednisone was started according to severity and topical clobetasol was tapered (Fig. 1). Nivolumab was restarted. Total IgE level was elevated and complete blood count revealed an increased absolute eosinophil count compared to pre-treatment levels, but it remained within normal ranges. The patient experienced a near-complete response to nivolumab, as shown by CT scan performed on August 2017. At this time, he presents only some lesions in the lower lip and he continues treatment with prednisone 5 mg bid and intermittent topical steroids.

Conclusion: Although we cannot be sure that BP is due to nivolumab, the timing of cancer diagnosis (24 months prior) compared with nivolumab initiation (12 months prior) and the fact that the other medications were tolerated for years without similar cutaneous lesions argue against a paraneoplastic BP or BP-related to another medication, respectively [1,2]. About treatment of BP during anti-PD-1/PD-L1, our research in the literature found a case report of BP during nivolumab with elevated IgE level and successfully treated with omalizumab, thus sparing steroids that could interfere with the antitumor activity of nivolumab [2,3]. Finally, all but few BP cases in the setting of anti-PD-1/PD-L1 therapy were associated with partial/complete response or stable like: is there a relationship between development of BP and anti-cancer responses [1,2]?

Consent for publication: The authors declare that written informed consent was obtained from the patients for publication.

Melanoma bridge 2017—Keynote speaker presentations

System biology session in melanoma

K4

Tumor and host genomics regulating immunotherapy efficacy

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Two major phenotypes of human melanoma metastases have been observed based on gene expression profiling and confirmatory assays. One subgroup of patients has a T cell-inflamed phenotype that includes expression of chemokines, T cell markers, and a type I IFN signature. In contrast, the other major subset lacks this phenotype and appears to display immune ‘exclusion’. The mechanisms of immune escape are likely distinct in these two subsets, and therefore the optimal immunotherapeutic interventions necessary to promote clinical responses may be different. The T cell-inflamed tumor microenvironment subset shows the highest expression of negative regulatory factors, including PD-L1, IDO, and FoxP3+ Tregs. Deep analysis of tumor antigen-specific T cells in the tumor microenvironment has identified additional mechanisms of immune dysfunction and new potential therapeutic targets. Treatment strategies targeting several pathways have been translated back into the clinic, including anti-PD-1/PD-L1 mAbs and IDO inhibitors, and combinations of these agents also look promising. In contrast to the T cell-inflamed melanomas, non-T cell-inflamed tumors are largely immunotherapy resistant with current approaches. Natural innate immune sensing of tumors appears to occur via the host STING pathway, type I IFN production, and cross-priming of T cells via CD8+ DCs, and these factors are absent in non-T cell-inflamed tumors. New strategies are being developed to engage or mimic this pathway as a therapeutic endeavor, including STING agonists. The molecular mechanisms that mediate the absence of the T cell-inflamed tumor microenvironment in patients are being elucidated using parallel genomics platforms. The first oncogene pathway identified that mediates immune exclusion is the Wnt/β-catenin pathway, which argues that new pharmacologic strategies to target this pathway should be developed to restore immune access to the tumor microenvironment. Recent evidence has indicated that host factors, including the intestinal microbiota, are also critical. We recently have identified commensal bacteria in mouse models that augment spontaneous anti-tumor immunity and increase efficacy of anti-PD-L1 therapy. Similar analyses in human melanoma patients have been performed, and commensal bacteria have similarly been identified that correlate with anti-PD-1 efficacy. These results have prompted the development of new probiotics as potential therapeutics, to improve spontaneous immune infiltration and expand immunotherapy efficacy in the clinic.
Melanoma bridge 2017—oral presentations

System biology session in melanoma

O14 Monitoring of melanoma clinical progression by circulating NK and T cells immunoprofiling: a potential role for CCR7 in metastatic spread
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Background: Natural Killer (NK) cells selectively recognize lymph node associated melanoma metastasis cells [1]. Peripheral blood frequencies of CD56bright and CD56dim NK cell subsets are subverted in stage III melanoma patients, while a high frequency of CCR7+KIR+CD57+CD56dim NK cells in melanoma colonized lymph nodes directly correlates with the patients survival [2]. Here, we validate and identify additional changes in the NK cells repertoire characterizing the transition from the different stages of melanoma that can improve the patients diagnosis, and dissected CCR7 potential role in metastatic process by measuring it on both melanoma and NK cells.

Results: Immuno-profile of 42 healthy donors and 103 melanoma patients (stage II, III, IV), together with biological variables, was used to create an OPLS-DA multivariate model. The model gave a good separation between healthy donors and the three groups of patients (Fig. 1A). Immuno-profile of both stage II and III melanoma patients showed an increased CXCR2 percentage, as previously observed [2], and a reduced CD57 frequency and NKp46 expression on the NKdim cells, that correlated with lack of responsiveness to K562 cells pulsing (Figure 2). Instead, stage IV melanoma patients showed high frequencies of CCR7+CD56bright NK cells, which displayed a longitudinal steady increase during the disease evolution.

Fig. 2 NK and T cells function analysis in healthy donors and mela-

Cytokine profile analysis showed a progressive sera accumulation of MCP-1, IL-6, IL-8 IL-15 and CCL19 (Table 1), with the first three that have been demonstrated to be produced by primary melanoma cells and found in TILN milieu [2], while IL-15 was involved in phenotypic changes of NK cells from melanoma patients [3]. CCL19 longitudinal increase over the clinical evolution, perfectly matched with the accumulation of circulating CD56brightCCCR7+ cells subset (Table 1).

Table 1 Univariate analysis of melanoma patients sera cytokines

| Cytokine | Physiological range (pg/ml) | Mean ± SD stage II (pg/ml) | Mean ± SD stage III (pg/ml) | Mean ± SD stage IV (pg/ml) | p value |
|----------|-----------------------------|-----------------------------|-----------------------------|-----------------------------|--------|
| MCP1     | 70.5–203                    | 232.1 ± 54.0                | 225.4 ± 69.1                | 222.9 ± 77.7                | 0.0085 |
| IL-6     | 0–5.6                       | 1.9 ± 2.2                   | 1.1 ± 1.0                   | 1.2 ± 1.4                   | <0.0001|
| IL-8     | 1.9–17.4                    | 9.1 ± 8.1                   | 13.7 ± 18.3                 | 496 ± 51.7                  | 0.0013 |
| IL-15    | 0–3.9                       | 2.0 ± 0.3                   | 1.7 ± 0.5                   | 4.0 ± 1.9                   | <0.0001|
| CCL19 N. T. | 164.9 ± 132.5                | 586.6 ± 73.18028 ± 4036     | 0.0490                      |

CCR7 analysis on melanoma cells showed that it is expressed by a small fraction (1–5%), which is further characterized by a selective co-expression between CCR7 and the inhibitory ligands PD-L1 and Galec- tin-9 (Fig. 3).

Fig. 3 PD-L1 and Tim-3 frequency and expression on CCR7— (white bars) and CCR7+ (Black bars) melanoma cells

Conclusions: Our data suggest a role of CCR7 expression on both melanoma and NK cells that can explain the lymph node tumor colonization and the conversion of the melanoma lesion from local to systemic.

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O15 Outcome prediction on melanoma patients subjected to immunotherapy treatments by 1H-NMR metabolomic profiling approach
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Fig. 4 Liver 1H-NMR metabolomic profiles of healthy donors and mel-

Fig. 4 Platelet metabolomics, 1H-NMR profiling, showed a dysregulation of the N-acetylaspartate (NAA) and creatine levels (black bars) and an increased levels of glutamate (grey bars) and glutamine (blue bars) in platelets of stage IV melanoma patients compared to healthy donors and stage II and III melanoma patients.

Conclusions: The 1H-NMR metabolomics results demonstrated a platelet metabolic signature that could be used for melanoma outcome prediction.

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Background: The evaluation of metabolomic profiling in biological fluids is recently emerged as a powerful and reliable tool for the identification of novel biomarkers to improve early diagnosis and prognosis classification, as well as prediction of treatment benefit in cancer patients. In this context, Nuclear Magnetic Resonance (NMR) spectroscopy represents the only non-destructive technique able to identify and quantify complex mixtures of metabolites by using small sample volumes and a fast sample preparation approach. In our laboratory we performed metabolomic profiling on liquid biopsy samples collected in eighty metastatic melanoma patients subjected to different immunotherapy treatments (i.e. anti-PD1 MoAb nivolumab and pembrolizumab, anti CTL4 MoAb ipilimumab).

Materials and methods: A 600 MHz Avance Bruker spectrometer with cryoprobe, equipped with an automation system was used to acquire 1H-NMR spectra on all samples at 300 K. The spectral 0.50–8.60 ppm region of the 1H-NMR spectra was integrated in buckets of 0.04 ppm using the AMIX package. The bucketed regions were normalized to the total spectrum area using Pareto scaling. Orthogonal Projections to Latent Structures Discriminant Analysis (OPLS-DA) and Loading Plot were used by the MetaboAnalyst tool to analyze and to compare the different groups.

Results: The patients were subdivided in two sub-groups to obtain a training set and a validation set. Hence, firstly 1H-NMR spectra were acquired and analyzed on samples belonging to training set. Through this procedure, we were able to group in separate clusters the patients with different outcome in terms of overall survival and to identify a set of metabolites that can discriminate patients before treatments with favorable than those with worst outcome. Then, we validated the significance of the selected metabolites in the validation set. Finally, ROC plots and Kaplan–Meier curves evidenced the metabolites correlated with overall survival.

Conclusion: Overall, these data suggest that metabolomic profiling performed on liquid biopsy samples by NMR is a potent and affordable method to improve outcome prediction for melanoma treatment.

O16 MiRNA analysis as a complement to screening procedures for early diagnosis of Malignant Melanoma (MM)
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Background: MM is the most aggressive skin cancer, its incidence doubled over the past 10 years and its mortality is still around 80%, in the advanced disease. The goal of the present study is to find new more effective screening methods, and to identify MM at high risk of recurrence [1, 2]. In more details, the aim of the project is to investigate the role of specific miRNAs, as a screening tool of MM [3-6].

Materials and methods: Individuals who joined the melanoma screening at the “Lega italiana per la lotta contro i tumori” (LILT), and with a suspect skin lesion, after signing informed consent, underwent a peripheral blood sample collection before the surgical removal. We compared the plasma level of 15 miRNAs in confirmed MM patients, versus non-confirmed. The study planned to enrol 700 subjects, to identify 100 MM.

The considered miRNAs (miR-15b, miR-122, miR-126, miR-150, miR-155, miR-182, miR-199, miR-200c, miR-205, miR-211, miR-23b, miR-33a, miR-34a, miR-424 and miR-29c) were selected on the basis of literature data and measured by RT-PCR.

Results: From September 2014 to December 2016, 16% (101/633) of enrolled individuals had histologic confirmation of MM: 38 in situ and S9 infiltrating, instead 4 patients were ineligible and were excluded from analysis; Among the patients without MM confirmation, we selected 100 controls who best matched the characteristics of the experimental population.

Till now we have analysed the role of miRNAs as potential tools for screening, while their prognostic role is not yet evaluated. The analysis showed a significant reduction in the expression of miR-199a-5p (p = 0.0027) and miR-122-5p (p = 0.05) in the MM (N = 97) group compared to the controls (N = 100). Multivariate analysis is underway in order to associate miRNA expression profiling with clinical-pathological characteristics.

Conclusions: The data available today do not allow to identify a miRNA that can be used as a complementary tool for the screening of early-stage MM. Significant reduction of miR-122-5p and miR199-5p in MM compared to controls, however, leads us to deepen their biological role in the pathogenesis of MM and their possible association with clinical-pathological features of subjects under screening. Multivariate analysis data will be available and presented at the meeting.

Acknowledgements: This project has been realized with the support of the CRC Foundation and the LILT section of Cuneo.

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O17 Predictive biomarkers of check point inhibition toxicity in metastatic melanoma
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Background: There are no predictive biomarkers of ipilimumab (IPI) toxicity. Of metastatic melanoma (MM) patients (pts) receiving IPI (3 mg/kg), 35% require systemic therapies to treat immune-related adverse events (irAEs) and 20% must terminate treatment [1]. Here we tested the hypothesis that a pre-existing autoantibody (autoAb) profile is predictive of IPI irAEs.

Methods: We measured autoAb levels in pre- and post-treatment sera from MM pts who received IPI (3 mg/kg) monotherapy on a prototype microarray containing ~ 20,000 unique full-length human proteins (HuProt array, CDI Laboratories). Clinical data were prospectively collected with protocol-driven follow-up. IraEs were categorized by CTCAE guidelines as none (grade 0), mild (grade 1–2), or severe (grade 3–4). AutoAb levels were standardized using median quantile normalization and considered positive hits if ≥ 2-SD above the peak array signal and differed by ≥ 2-fold with p < 0.05 between toxicity groups (Non-parametric Analysis/Wilcoxon test).
**Results:** Seventy-eight sera from 37 MM pts were analyzed. Antibodies against CTLA-4 were significantly elevated post IPI treatment (p < 0.0001), validating the assay. The pre-treatment levels of 190 IgG autoAbs were significantly different in pts who experienced iAEs (n = 28) compared to those with no iAEs (n = 9). Comparison of severe iAEs (n = 9) and no iAEs (n = 9) groups revealed 128 IgG autoAbs that significantly differed in pre-treatment sera. Localization and pathway analysis (UniProt, KEGG, Reactome) showed 81/190 (43%) of the autoAbs targeted nuclear and mitochondrial antigens and were enriched in metabolic pathways (p = 0.015). AutoAbs associated with iAEs did not correlate with treatment response.

**Conclusions:** AutoAbs to antigens enriched in metabolic pathways prior to treatment may predict IPI-induced toxicity in MM. The subcellular localization of targeted antigens could explain the autoimmune toxicities associated with IPI. Studies in larger cohorts and in pts receiving other checkpoint inhibitors and/or combination therapies are essential to determine the validity of the data. If validated, our results would support the discovery of the first toxicity predictor in cancer immunotherapy.

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**O18 Soluble CD73 in the peripheral blood: a potential biomarker in patients with advanced melanoma receiving nivolumab**

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**Journal of Translational Medicine 2018, 16 (Suppl 1):O18**

**Background:** Anti-PD1 agents are successfully used in therapy to treat patients with advanced melanoma. Here, we retrospectively analysed the CD73 enzyme activity in the peripheral blood of patients with metastatic melanoma receiving nivolumab. CD73 is an ectonucleotidase able to generate adenosine from AMP. Adenosine in the tumor microenvironment is a potent immune-suppressive mediator, so that inhibition of CD73-generating enzyme or blockade of adenosine receptors is a promising therapeutic strategy to fight cancer.

**Materials and methods:** CD73 enzyme activity was retrospectively analysed in the plasma of patients before receiving nivolumab. CD73 activity was measured in frozen peripheral blood mononuclear cells (PBMCs) that were thawed and then rested briefly, and subjected to flow cytometry analysis for myeloid-derived suppressor cells (MDSCs: CD14+ /CD33+ /CD11b+ /HLA-DR-/low), CD8+ and CD4+, alone or in association with PD-1 and CD73+.

**Results:** Our data demonstrated that patients with lower baseline values of CD8+/PD-1+ /CD73+ had high OS and PFS (34.8 and 19.2 months, respectively); conversely, patients with higher baseline frequency of these cells experienced lower OS and PFS (9.5 and 2.8 months, respectively; OS p < 0.003, PFS p < 0.007) (Tables 1, 2). In addition, increasing number of total CD8+ cells (p < 0.05) and especially of CD8+/PD-1+ cells (p < 0.04) were negatively correlated with survival (Table 1).

**Table 1 Median OS (months) for patients with values under and over the median of CD8+, CD8+/PD-1+ and CD8+/PD-1+/CD73+**

| Factors (median) | Median OS (months) for patients with values under the median | Median OS (months) for patients with values over the median | p value |
|-----------------|-------------------------------------------------------------|-------------------------------------------------------------|---------|
| CD8+ (250)      | 204                                                        | 69                                                         | 0.05    |
| CD8+/PD-1+ (7.8)| 34.8                                                       | 9.9                                                         | 0.04    |
| CD8+/PD-1+/CD73+ (2.3) | 34.8                                                      | 9.5                                                         | 0.003   |

We also postulate that increased levels of CD73 may contribute to affect the response of immunotherapeutic agents in cancer patients.

**Consent to publish:** All the patients provided written informed consent.

**O19 Expression of CD73 on CD8+/PD-1+ cells as a new possible biomarker for advanced melanoma patients treated with nivolumab**

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**Background:** Anti-programmed death (PD)-1 monoclonal antibodies have changed the prognosis of metastatic melanoma, improving overall survival [1]. However, still a proportion of patients is unresponsive to these compounds, indicating the presence of other immunosuppressive mechanisms. Thus, the identification of reliable biomarkers to predict the response to checkpoint blockades is still an unmet need.

Recent findings showed a tumor-induced immunosuppressive pathway, in which the extracellular adenosine produced by tumor-derived enzyme CD73 (5’-ectonucleotidase) promotes tumor growth by inhibiting immunosuppressive T-cell action [2]. Targeting adenosine generation by blockade of CD73 significantly enhances anti-tumor activity of anti-PD-1 drugs, inducing full regression in some tumor models [3]. The aim of the study was to investigate whether baseline levels of CD73+ on circulating CD8+, CD4+ and MDSCs cells could be considered as potential biomarkers in stage IV melanoma patients treated with nivolumab.

**Materials and methods:** Blood samples from 36 advanced melano-ma patients were taken before nivolumab treatment; blood popula-tions were measured in frozen peripheral blood mononuclear cells (PBMCs) that were thawed and then rested briefly, and subjected to flow cytometry analysis for myeloid-derived suppressor cells (MDSCs: CD14+ /CD33+ /CD11b+ /HLA-DR-/low), CD8+ and CD4+, alone or in association with PD-1 and CD73+.

**Results:** The aim of the study was to investigate whether baseline levels of CD73+ on circulating CD8+, CD4+ and MDSCs cells could be considered as potential biomarkers in stage IV melanoma patients treated with nivolumab. The aim of the study was to investigate whether baseline levels of CD73+ on circulating CD8+, CD4+ and MDSCs cells could be considered as potential biomarkers in stage IV melanoma patients treated with nivolumab.
Table 2 Median PFS (months) for patients with values under and over the median of CD8+/PD-1+/CD73+

| Factors (median) | Median PFS (months) for patients with values under the median | Median PFS (months) for patients with values over the median | p value |
|------------------|-------------------------------------------------------------|-------------------------------------------------------------|---------|
| CD8+/PD-1+/CD73+  | 19.2                                                        | 2.8                                                         | 0.007   |

Furthermore, the baseline values of MDSCs/CD73+ and CD4+/CD73+ cells showed no significant differences in survival.

**Conclusions:** Our work indicates that the analysis of CD8+/PD-1+/CD73+ baseline levels in advanced melanoma patients treated with nivolumab could be associate to treatment outcome. Also, these preliminary results suggest that the therapeutic potential of anti-CD73 inhibitors, which are in phase I of clinical trials, increasing the development of new treatment combinations strategies with other immune checkpoint monoclonal antibodies. Further studies on a larger number of patients are ongoing to confirm the data obtained.

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**Combination strategy session**

**O20**
**Prognostic impact of baseline tumor immune infiltrate on disease-free survival (DFS) in patients with completely resected, BRAF+/BRAF V600E+ melanoma receiving adjuvant vemurafenib**

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**J Transl Med 2018, 16 (Suppl 1):O20**

**Background:** The BRIM8 study (clinicaltrials.gov NCT01667419) evaluated the efficacy of adjuvant vemurafenib in patients with fully resected BRAF+/BRAF V600E+ melanoma at high risk of recurrence.

**Materials and methods:** This 2-cohort (C) study randomized 498 adults with resected stage IIC, IIIA, or IIIB melanoma (C1) or stage IIIC melanoma (C2) 1:1 to vemurafenib 960 mg BID or placebo for 52 weeks. The primary endpoint was DFS. Hierarchical analysis of C2 before C1 was prespecified. The association of tumor immune infiltrate with DFS was explored retrospectively.

**Results:** In C2, median DFS was greater with adjuvant vemurafenib vs placebo (23.1 vs 15.4 mos), but the risk reduction was not statistically significant (log-rank p = 0.2598). Vemurafenib reduced DFS risk by 46% vs placebo in C1 (median not estimable vs 36.9 mos; log-rank p = 0.0010, not statistically significant as the primary DFS endpoint was not met in C2). Tumor samples were available for ~60% of patients. In the pooled biomarker population, placebo-treated patients with < 1% CD8+ T cells in the tumor center had shorter median DFS vs patients with ≥ 1% CD8+ T cells (7.7 vs 47.8 months). The DFS benefit from vemurafenib vs placebo was greater in patients with < 1% CD8+ T cells in the tumor center (hazard ratio [HR] 0.56, 95% CI 0.34–0.92) than in patients with ≥ 1% CD8+ T cells (HR 0.77, 95% CI 0.48–1.22). Likewise, placebo-treated patients with < 5% PD-L1+ immune cells (IC) had shorter median DFS vs patients with ≥ 5% PD-L1+ IC (7.2 vs 47.8 months). The DFS benefit from vemurafenib vs placebo was greater in patients with < 5% PD-L1+ IC in their tumors (HR 0.36, 95% CI 0.24–0.56) than in patients with ≥ 5% PD-L1+ IC (HR 0.99, 95% CI 0.58–1.69). The presence of CD8+ T cells and PD-L1+ IC are favorable prognostic factors for DFS.

**Conclusion:** Treatment with adjuvant vemurafenib may overcome the poor DFS prognosis associated with low CD8/PD-L1 expression in the tumor.

**O21**
**4SC-202 plus anti-PD1: breaking PD1-refractoriness to increase efficacy of checkpoint inhibition in patients with advanced melanoma**

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**Journal of Translational Medicine 2018, 16 (Suppl 1):O21**

**Background:** Despite successes in the treatment of melanoma patients with checkpoint inhibitors (anti-PD1 antibodies), the majority of patients do not respond to checkpoint inhibition alone and a high unmet medical need remains for these patients. One promising approach is to increase the number of patients benefiting from checkpoint inhibition by enhancing the immunogenicity and alter the tumor microenvironment from a more immune-deserted to an immune-inflamed phenotype by means of combination therapy. Epi-genetic modulation has been reported as one key determining factor in shaping the immune microenvironment and compounds altering these processes [e.g. histone deacetylases (HDAC) inhibitors] are particularly promising.

**Results:** Here, we report results for 4SC-202, an orally available clinical stage HDAC inhibitor, and outline the further clinical development. 4SC-202 treatment led to an increase of MHC class II molecules and enhanced expression of inflammatory markers like IFN-γ and various chemokines in tumors. Furthermore, detailed analysis of the tumor microenvironment in tumor-bearing animals revealed that 4SC-202 strongly altered the immune cell composition and particularly the number of cytotoxic T cells (CTL) was markedly increased. Importantly, subsequent combination treatment of 4SC-202 with checkpoint inhibitors in syngenic animal models showed a strong synergistic effect resulting in an increased tumor growth reduction.

For the further clinical development, start of a Phase Ib/II clinical study (‘SENSITIZE’; ClinicalTrials.gov Identifier: NCT03278661) is planned by the end of this year. This study, conducted in Germany with up to 6 sites (~30 patients) will enroll patients with advanced cutaneous melanoma who are refractory/non-responding to treatment with anti-PD-1 antibodies. These patients clearly represent a population with a high unmet medical need and might be characterized by an unfavorable tumor immunology and microenvironment for immunotherapy in general and checkpoint inhibition in particular.
Conclusion: We hypothesize that addition of 4SC-202 to anti-PD-1 antibody treatment may lead to increased immunogenicity of the tumor, an inflamed tumor microenvironment and ultimately to clinical benefit in anti-PD-1 refractory/non-responding advanced melanoma patients.

Melanoma bridge 2017—poster presentations

P4 Insights into the patient perspective: uncovering the clinical experiences and unmet needs for melanoma patients

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Background: Data from the literature indicate that the perception of patients with melanoma regarding the support received from healthcare services is generally inadequate with respect to the patients' need. Approximately 30% of melanoma patients face significant physical and psychological issues related to the diagnosis, treatment, and follow-up [1-3]. The aim of this study is to evaluate the patients’ specific clinical and communication needs in order to maximize their quality of health care.

Materials and methods: A predefined questionnaire was submitted to 130 patients with new diagnoses of melanoma (stage I-IV), followed at San Raffaele Hospital (HSR) from January 2015 to January 2017. The study was approved by the Ethical Committee. The participants were recruited using non-probability purposive sampling techniques. The questionnaires were administered by email or telephone. It included 34 items, evaluating patients’ demographics, education and employment status, basic knowledge on melanoma, and the understanding of diagnosis, stage, treatment details, prognosis, screening behaviors, quality of medical-patient communication and the perception for healthcare personnel support. The questionnaires were adapted from those used in Australia and New Zealand [4–5]. The reliability of the instrument was tested using test–retest reliability in IBM SPSS version 18. Statistical analysis was carried out using excel and PASW version 18 (SPSS, Inc, II), adjusted by age, gender, education and stage of tumor.

Results: One hundred and thirty patients accessed the questionnaire. Fifty-two have been already completed. Here we present preliminary date. In Table 1 patients’ characteristics.

Table 1 Patients’ characteristics

| Patients’ characteristics | Total no 52 (100%) |
|---------------------------|-------------------|
| Gender                    |                   |
| Female                    | 33 (63.5%)        |
| Male                      | 19 (36.5%)        |
| Education level           |                   |
| < high school             | 8 (15.4%)         |
| High school               | 8 (15.4%)         |
| Apprenticeship training   | 21 (40.4%)        |
| Bachelors/masters         | 3 (5.8%)          |
| Doctorate                 | 12 (23.1%)        |
| Tumour’ sites             |                   |
| Torso                     | 25 (48.1%)        |

Reference

1. Living as a melanoma skin cancer survivor. American Cancer Society, https://www.cancer.org/cancer/melanoma-skin-cancer/after-treatment/follow-up.html.

P5 Oncolytic immunotherapy with PV-10

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Background: PV-10 is an investigational small molecule oncolytic immunotherapy (10% w/v rose bengal disodium for injection). Intradermal (IL) injection can elicit a primary, local effect in injected tumor tissue (immunogenic cell death and release of tumor antigens) capable of stimulating a secondary, functional systemic immune response (including tumor-specific reactivity in circulating T cells in treatment-refractory patients). PV-10 has been investigated in Phase 1 and 2 clinical trials in 130 patients with melanoma; an expanded access protocol accrued an additional 180 melanoma patients.

Materials and methods: PV-10 is currently under clinical investigation in two melanoma studies. Phase 3 study PV-10-MM-31 (NCT02288897), an international multicenter, randomized controlled trial (RCT), is

Table 1 continued

| Patients’ characteristics | Total no 52 (100%) |
|---------------------------|-------------------|
| Limbs                     | 19 (36.5%)        |
| Head or neck              | 5 (9.6%)          |
| Others                    | 3 (5.8%)          |
| Tumour’ stage             |                   |
| Stage I                   | 29 (55.8%)        |
| Stage II                  | 4 (7.7%)          |
| Stage III                 | 3 (5.8%)          |
| Stage IV                  | 4 (7.7%)          |
| Unknown to the patient    | 12 (23.1%)        |

Patients were grouped according to age (< 50, 50–70 and > 70 years) and the level of education (high school or less and vocational training). Sixty-three per cent of patients were satisfied about the support received, and 51.4% received enough information regarding diagnosis, prevention, treatment options and recurrence of the disease. Patients underline that like to have a single healthcare consultant.

Conclusions: This is a deepened patient-centred approach conceived for melanoma patients that would provide a perspective on melanoma care, highlighting the areas that require the definition of a new protocol based on patients and their caregivers’ needs. In Table 1 patients’ characteristics.

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1. Living as a melanoma skin cancer survivor. American Cancer Society, https://www.cancer.org/cancer/melanoma-skin-cancer/after-treatment/follow-up.html.

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assessing effectiveness of PV-10 compared to the investigator’s choice of systemic chemotherapy or intralesional oncolytic viral therapy in up to 225 patients with locally advanced cutaneous melanoma (AJCC Stage IIIb to IV M1a). The primary endpoint is progression-free survival (PFS) assessed via RECIST 1.1. Phase 1b/2 combination study PV-10-MM-1201 (NCT02255732), an international multicenter, open-label, sequential phase study, is assessing safety and efficacy of PV-10 in combination with systemic immune checkpoint inhibition. Stage IV metastatic melanoma patients with at least one injectable cutaneous or subcutaneous lesion who are candidates for pembrolizumab are eligible for study participation. In the Phase 1b portion of the study, up to 24 subjects will receive the combination (i.e., PV-10 + SoC) with PFS and objective response (OR) as key secondary endpoints. In the Phase 2 portion of the study up to 120 study participants will be randomized 1:1 to receive either the combination or pembrolizumab alone (i.e., PV-10 + SoC vs SoC) with PFS as the primary endpoint.

Results: Both studies are currently enrolling patients. The Phase 3 study is open at centers in the USA, Australia, Italy and Germany; additional regulatory approval has been received in Mexico, with approval pending in France, Poland and Argentina. The Phase 1b study is open at centers in the USA and AUS. To date, no unexpected safety signals have been observed in either study, and Phase 2 of the combination study is expected to begin in early 2018. Preliminary efficacy data from Phase 1b of the combination trial-in-progress will be presented.

Conclusion: Pivotal testing of single agent PV-10 is ongoing. Conclusion of Phase 1b testing of PV-10 in combination with checkpoint inhibition is anticipated to lead to Phase 2 randomized testing in 2018.

**P6 The reciprocal interplay between miR-579-3p and MITF controls melanoma proliferation and resistance to target therapy**
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**Background:** MAPK signaling is the main oncogenic driver in metastatic melanomas bearing mutations in BRAF kinase. These tumors are currently treated with the combination of BRAF/MEK inhibitors (MAPKi), but this therapy is plagued by drug resistance. In this context we recently focused on non-mutational mechanisms contributing to the development of drug resistance and in particular on the role of miRNAs. This led us to the discovery of miR-579-3p as an antagonist of melanoma progression and resistance. miR-579-3p targets both BRAF and MDM2 oncogenes and its expression is strongly downregulated during melanoma development and during progression to drug resistance. In the present work, we studied the mechanisms regulating the expression of miR-579-3p and the interplay with MITF, the master regulator of the melanocyte lineage.

**Materials and methods:** JASPAR open database was queried to identify potential transcription factors binding to the miR-579-3p promoter, which were confirmed through silencing, luciferase and Chromatin-Immunoprecipitation experiments. Three melanoma cell lines (M14, LOX IMVI and WM266) bearing different mutations in the BRAF oncogene were exposed to MAPKi for different times and then collected for protein and RNA extraction. FFPE samples from matched tumors before and after the development of resistance to MAPKi were used for total RNA extraction and analysis through qRT-PCR.

**Results:** miR-579-3p is located within intron 11 of the Zinc Finger Recombinase (ZFR) gene. Bioinformatic analysis allowed us to identify two putative MITF binding sites within ZFR gene promoter. Gene silencing, luciferase assays and ChIP experiments confirmed that MITF binds to the ZFR promoter and that acts a positive regulator of miR-579-3p transcription. Moreover, miR-579-3p by targeting BRAF is able to stabilize MITF protein thus inducing its own transcription in a positive feedback regulatory loop. In line with these findings, BRAF mutated melanoma cells exposed to MAPKi inhibitors undergo a rapid upregulation of both MITF and miR-579-3p which corresponds to cell growth inhibition. On the opposite drug resistant melanoma cells show downregulation of MITF and miR-579-3p in parallel. Simultaneous loss of both MITF and miR-579-3p is also observed in tumor samples from patients after development of drug resistance.

**Conclusions:** In this work we uncover a positive feedback regulatory loop involving MITF and the on co-suppressor miRNA miR-579-3p which inhibits melanoma proliferation. Drug resistant melanoma cells escape this mechanism by downregulating both MITF and miR-579-3p. We are currently investigating the molecular basis of the "switch" from MITF/miR-579-3p+ drug sensitive to the MITF/miR-579-3p- drug insensitive status.

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