Fluctuation of systemic immunity in melanoma and implications for timing of therapy


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1. ABSTRACT

Evidence suggests that immunological response in chronic inflammation is dynamic, oscillating between active immunity and tolerance. We hypothesized that a similar dynamic exists in melanoma and administration of therapy during a unique phase of such oscillation could impact clinical outcome. Patients with metastatic melanoma eligible to undergo temozolomide underwent serial measurements of C-reactive protein (CRP) and immune biomarkers every 2-3 days for 2 weeks before starting therapy. Treatment was initiated prior to the estimated next CRP peak, or on day 14 post-registration if a peak was not identified. Time profiles of measured biomarkers were analyzed by fitting serially measured data points to 9 mathematical functions and were correlated to time of therapy and outcome. Data suggested that metastatic melanoma patients exhibit a dynamic immune response. The fluctuation of several biomarkers fitted cosine functions with periods which were multiples of 3-4 days. Chemotherapy delivery during a unique phase of this cycle seemed to correlate with improved response. Individualized conventional chemotherapy delivery by synchronizing treatment with pre-existing patient-specific biorhythms may improve clinical outcomes in metastatic melanoma.

2. INTRODUCTION

The incidence of melanoma has been increasing over the last two decades at a rate higher than most malignancies (1). It is currently estimated that one in every 40 people in the United States will be diagnosed with melanoma during their lifetime (1, 2). While potentially curable when diagnosed early, the median survival of patients with metastatic disease remains less than 1 year, and less than 5% of patients are alive at 5 years (3). Melanoma is notoriously refractory to most chemotherapeutic agents, but does show sporadic and dramatic responsiveness to immune modulating therapies. One explanation for the isolated successes of immunotherapeutics is the increasingly recognized variability in systemic immune dysfunction in patients with disseminated disease. The abnormalities of immune competence in stage IV melanoma have been extensively studied; recent evidence suggests the existence of varying degrees of immune suppression in the tumor microenvironment, as well as systemic immune “exhaustion” brought upon by tumor-driven chronic inflammation (systemic helper T cells, Th2 dominance) (4). Different levels of such immune compromise in un-selected patients with stage IV melanoma may render different levels of “responsiveness” to immune stimulating drugs.
Many approaches have been used to generate endogenous anti-tumor immunity in metastatic melanoma (5-7). Recently, adoptive transfer studies have definitively demonstrated that some form of “re-setting” (lymphodepleting) endogenous immune homeostasis may be necessary for the success of infused, in vitro prepared, tumor specific cytotoxic T cells (8, 9). This approach implies that the state of cancer-associated immune dysfunction is established and static, requiring depletion. However, increasing pre-clinical data suggest that immunity to cancer is a cyclical process, which oscillates between active immune responsiveness and tolerance (10-12). This “on-off” cycle of systemic anti-tumor immunity is modulated, among others, by periodical clonal expansion of regulatory T-cells (Treg) and possibly other immunosuppressive elements (13-15). In animal models, Treg appear to undergo synchronous clonal expansion approximately 10 to 14 days post viral or tumor inoculation (16) and act to suppress the activity of effector T cells undergoing clonal expansion some days prior (13,17). Application of a single dose of cytotoxic (lymphodepleting) chemotherapy at the time when Treg enter mitosis and are undergoing clonal expansion postulated that delivery of TMZ just prior to the presumed time during said cycle to deliver TMZ (10,19). We demonstrated that some form of “re-setting” unknown pre-existing endogenous immunity may not be sufficient to achieve clinical benefit.

In the current study, we hypothesized that systemic (homeostatic) immunity in patients with metastatic melanoma exists in a state of dynamic equilibrium and that delivery of lymphodepleting cytotoxic chemotherapy (temozolomide, TMZ) in a unique phase of this cycle may favor active immunity over tolerance, yielding clinical benefit. As serum C-reactive protein (CRP) levels appeared to inversely correlate with regulatory T cell (Treg) frequency in patients with chronic inflammatory diseases (19), we used serial CRP measurements as a surrogate for Treg frequency to detect oscillations of systemic immunity and empirically select a time during said cycle to deliver TMZ (10,19). We postulated that delivery of TMZ just prior to the presumed peak of Treg concentration in the blood may yield maximum Treg depletion, favoring persistence of active systemic immunity.. The study objectives were: (1) ascertain whether or not CRP and other biomarkers of inflammation) is predictably variable over time in patients with advanced melanoma (i.e. is systemic immune competence is a dynamic process oscillating between states of “active” immunity and tolerance); and (2) therapeutically utilize this information and devise a treatment strategy to selectively lymphodeplete the immune down-regulating phase of systemic immunity.

3. MATERIALS AND METHODS

3.1. Patient population and study design

Eligible patients had unresectable, histologically confirmed stage IV disease, age over 18 years, measurable disease as defined by the Response Evaluation Criteria in Solid Tumors (RECIST), Eastern Cooperative Oncology Group (ECOG) performance status (PS) of 0-2, and life expectancy ≥ 3 months. Both previously untreated patients and patients who have had prior therapy for their metastatic disease (excluding prior exposure to TMZ) were eligible. All patients provided signed informed written consent, and the study was approved by the Mayo Clinic Rochester Institutional Review Board. The TMZ dose was 150 mg/m² on days 1-5 of cycle 1 and was increased to 200 mg/m² for all subsequent cycles if well tolerated. Patients were treated every 28 days until progression, unacceptable toxicity or patient refusal. Prior to initiation of first chemotherapy cycle, eligible patients underwent serial peripheral blood testing for CRP and associated immunological biomarkers every 2-3 days (to account for weekend days) for a period of two weeks. CRP concentration was measured using a clinical laboratory test (20) in real time in order to identify the optimal time for chemotherapy delivery. The time series of serial CRP concentration measurements was fitted to a sine curve and the next peak of CRP concentration was predicted based on the periodicity of CRP oscillations. Treatment with TMZ was initiated immediately prior to the estimated time of the next CPR peak, or on day 14 post-registration if the peak could not be identified.

3.2. Immunological studies

In order to study the global behavior of the anti-tumor immune response in metastatic melanoma, peripheral blood samples obtained prior to initiation of TMZ therapy were subsequently analyzed for plasma concentration of 29 different cytokines and 22 immune cell subsets (described below). All biospecimens were collected, processed, and stored following established and validated standard operating procedures in our laboratory (4). To reduce inter-assay variability, all assays were batchalyzed after study completion. All blood samples were collected at approximately the same time of day (between 0800h and 0100 h ) in order to minimize the contribution of circadian variation to the fluctuation of immune parameters.

3.2.1. Peripheral blood mononuclear cell (PBMC) immunophenotyping for immune cell subsets

Blood was separated into platelet poor plasma and PBMC using a density gradient (Ficol-hypaque, Amersham, Uppsala, Sweden). Plasma samples were stored at -70 °C, and PBMC were stored in liquid nitrogen. Immunophenotyping of PBMC was performed by flow cytometry using FITC- and PE-conjugated antibodies to CD3, CD4, CD8, CD16, CD56, CD62L, CD69, TIM3 (T-cell immunoglobulin domain and mucin domain 3), CD294,
Serially measured peripheral blood biomarkers were analyzed using CurveExpert 1.4 software (Daniel G. Hyams, Hixson, TN) and GraphPad Prism 4.0 software (GraphPad Software Inc. La Jolla CA) to construct time-dependent profiles of plasma cytokine/immune cell counts by fitting data points to selected mathematical functions. Technical reproducibility was assessed by the coefficient of variation (CV) among duplicates (average CV was 5.13% for 1593 data points). We used the correlation coefficient calculated by CurveExpert as the first criterion for goodness-of-fit, where S, considers the distribution around a constant line and is calculated as (\text{C} - \text{F})^2. S - \sum_{i=1}^{n} (\text{C}_i - \text{F}_i)^2 considers the deviation from the fitting curve and is calculated as . Using GraphPad S = \sum_{i=1}^{n} (\text{C}_i - \text{F}_i)^2 Prizzm we obtained R^2 values, 95% confidence intervals for the parameters of the fitted functions, and 95% confidence bands for the fitted curves. R^2 is calculated as R^2 = 1 - (S/S_0). These parameters were used as selection criteria in different steps of the analysis as described in the results section.

Most patients discontinued testing once the required number of data points needed for sine curve fitting and prediction of the CRP peak was obtained. Consequently, we extrapolated the value for cytokines/immune cell counts on the day of treatment. We also calculated the first derivative (FD) of the fitted function; this shows whether the function increases (positive value), decreases (negative value) or is steady (zero), and the magnitude of FD reflects the magnitude of the trend. The range of plasma cytokine concentrations and cell counts varied significantly among study subjects and they were converted into relative values with the following formula: (C_{max} - C_{min})/(C_{max} - C_{min}), where C_{max} is the maximum concentration within the observed time period, C_{min} is the minimum concentration within the same period, and C_{0} is the extrapolated concentration on the day of treatment. The same conversion was applied to FD values. In cases in which both maximum and minimum FD were negative we applied the formula: \text{FD} = -1*(\text{D}_{\text{max}} - \text{D}_{\text{min}})/\text{D}_{\text{max}} - \text{D}_{\text{min}}). Where D_{\text{max}} = \text{maximum FD within the observed time period}, D_{\text{min}} = \text{minimum FD within the same period}, D_{\text{a}} = \text{FD of the function for the extrapolated point corresponding to the day of treatment, in order to compensate for the subtraction of two negative numbers. Secondary data analysis and visualisation was done using Partek GS 6.5 software (Partek Inc. Saint Louis, MO).
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Table 1. Clinical and demographic characteristics of the 11 patients with metastatic melanoma enrolled in the study

<table>
<thead>
<tr>
<th>Patient #</th>
<th>Sex/ Age</th>
<th>Original diagnosis/ Anatomic site</th>
<th>Metastatic Disease</th>
<th>Nr. of prior treatments</th>
<th>Treatment start (# days after enrollment)</th>
<th>Nr. of TMZ cycles on study</th>
<th>PFS (days)</th>
</tr>
</thead>
<tbody>
<tr>
<td>#1</td>
<td>M/67</td>
<td>Nodular melanoma / ear</td>
<td>Lung Paratracheal LN</td>
<td>0</td>
<td>18</td>
<td>35</td>
<td>916</td>
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<tr>
<td>#2</td>
<td>M/55</td>
<td>Desmoplastic melanoma/scalp</td>
<td>Lung; liver; spleen; pancreas bone</td>
<td>0</td>
<td>12</td>
<td>1</td>
<td>37</td>
</tr>
<tr>
<td>#3</td>
<td>M/42</td>
<td>Nodular melanoma / limb</td>
<td>Subcutaneous limb/ trunk; adrenal glands bone</td>
<td>0</td>
<td>7</td>
<td>2</td>
<td>74</td>
</tr>
<tr>
<td>#4</td>
<td>M/82</td>
<td>Nodular melanoma / scalp</td>
<td>Soft tissue scalp/ neck/ upper limbs</td>
<td>0</td>
<td>11</td>
<td>24</td>
<td>841</td>
</tr>
<tr>
<td>#5</td>
<td>M/52</td>
<td>Cutaneous melanoma / scalp</td>
<td>Liver; Retropitoneal LN</td>
<td>2</td>
<td>14</td>
<td>2</td>
<td>91</td>
</tr>
<tr>
<td>#6</td>
<td>F/71</td>
<td>Cutaneous melanoma / scalp</td>
<td>Lung</td>
<td>1</td>
<td>17</td>
<td>4</td>
<td>132</td>
</tr>
<tr>
<td>#7</td>
<td>M/71</td>
<td>Cutaneous melanoma / limb</td>
<td>Lung; liver; pelvic LN</td>
<td>3</td>
<td>14</td>
<td>2</td>
<td>68</td>
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<tr>
<td>#8</td>
<td>M/51</td>
<td>Choroidal melanoma</td>
<td>Liver; bone</td>
<td>0</td>
<td>18</td>
<td>2</td>
<td>71</td>
</tr>
<tr>
<td>#9</td>
<td>F/72</td>
<td>Choroidal melanoma</td>
<td>Liver; spleen</td>
<td>0</td>
<td>20</td>
<td>2</td>
<td>74</td>
</tr>
<tr>
<td>#10</td>
<td>F/51</td>
<td>Cutaneous melanoma / scalp</td>
<td>Lung; soft tissue abdominal wall</td>
<td>2</td>
<td>15</td>
<td>2</td>
<td>70</td>
</tr>
<tr>
<td>#11</td>
<td>F/56</td>
<td>Acral melanoma</td>
<td>Lung; liver; soft tissue trunk; bone</td>
<td>4</td>
<td>2</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>#12</td>
<td>F/72</td>
<td>Amelanotic melanoma / limb</td>
<td>Lung; liver; soft tissue trunk; mesenterium; mediastinal &amp; retroperitoneal LN</td>
<td>0</td>
<td>20</td>
<td>2</td>
<td>77</td>
</tr>
</tbody>
</table>

1 Refers to prior chemotherapy, biochemotherapy (IL-2, IFN), GM-CSF, or melanoma vaccine. 2 Patient required hospitalization for complications secondary to metastatic disease soon after enrollment and did not receive any further therapy. 3 Insufficient number of peripheral blood samples obtained prior to initiation of treatment.

Figure 1. Time-dependent profiles of plasma cytokines and immune cell subsets fitted to 9 different mathematical models (functions) in 10 patients with metastatic melanoma. Function codes: F0= No Fit/No data, F1= Linear function y=ax+b; F2= Exponential function: y=ae^(bx); F3= Exponential Association: y=a(1-exp(-bx)); F4= Logistic function: y=a/(1+b*exp(-cx)); F5= Quadratic function: y=a+bx+cx^2; F6= Cosine function: y=a+b*cos(cx+d); F7= Rational function: y=(a+bx)/(1+cx+dx^2); F8= Gaussian function: y=a*exp(-(b-x)^2)/(2*c^2)); F9= MMF Model: y=(a*b+c*x^d)/(b+x^d); The goodness-of-fit was estimated using the correlation coefficient (CC) calculated by CurveExpert 1.4 software. The frequency distribution of the correlation coefficient was computed across all profiles and all patients and the value of the 75th percentile (0.86) was accepted as a cut-off to eliminate profiles which did not fit a model well. Black boxes represent the range of values from 25th to 75th percentiles and whiskers represent the range from 10th to 90th percentiles.
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Figure 2. Flowchart of data selection process. Time-dependent patterns and dependencies satisfying selection criteria were identified. The flowchart demonstrates the filtering algorithm used for data analysis and the results obtained at each step of this analysis with references to the figures in the manuscript.

was 66 years (range 42 to 82 years). The median number of TMZ cycles administered was 2 (range 1 to 35). Two complete tumor responses were documented; both patients have since discontinued the study drug after 35 and 24 cycles respectively, and remain disease-free at 39 and 31.5 months since study entry. All of the remaining 9 patients have discontinued treatment due to tumor progression and have subsequently died. Overall, the median progression-free survival (PFS) time and the median overall survival (OS) times were 74 days and 12.5 months, respectively.

4.2. Dynamic changes of systemic immunity in metastatic melanoma

We analyzed the time-dependent profiles of 51 longitudinally measured immune biomarkers (see immunological studies above) prior to initiation of TMZ therapy. Ten of the 12 patients enrolled in the study were studied; one patient canceled participation in the trial prior to starting TMZ therapy and another patient had an insufficient number of peripheral blood samples obtained prior to initiation of treatment. Serially measured data points were fitted to mathematical functions using previously described algorithms (21,22). Curve-fitting was performed based on 6 to 7 sequential measurements (time points) for each variable and each patient over a period of 14 days. The distribution frequencies of the 9 different mathematical models employed in the analysis showed that most time-dependent profiles fitted sinusoidal or rational functions implying that these immune parameters oscillate repeatedly in an apparent predictable fashion (Figure 1).

Next, we evaluated whether an ordered versus disordered pattern of fluctuation correlated with clinical outcome. In order to reveal this trend we applied a set of selection criteria aimed at filtering out insignificant dependencies (associations) (Figure 2). The criteria were set to verify that: i) experimental data fit a mathematical model (CC ≥ 0.86); ii) an ordered pattern is consistent across patients (indexing and ranking described below); iii) each of the 9 mathematical models is consistent across patients (same as (ii), but only for one specific model). Each curve fitting attempt was assigned an index of fitness. The index was 1 if the experimental profile fitted a model function well (CC ≥ 0.86) and the function was biologically possible. Functions with infinite growth or infinite decline were considered biologically implausible as their extrapolation produces biologically impossible values (e.g. < 0) for cytokine concentrations or cell counts, and they were assigned an index of zero (0). Finally the index was -1 if a profile did not fit any function. The sum of indices
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Table 2. The rank of measured immune biomarkers based on the likelihood of fitting a mathematical function

<table>
<thead>
<tr>
<th>Variable</th>
<th>Rank</th>
<th>Variable</th>
<th>Rank</th>
</tr>
</thead>
<tbody>
<tr>
<td>IL-10</td>
<td>7</td>
<td>CD3-/CD16-</td>
<td>3</td>
</tr>
<tr>
<td>IL-12p70</td>
<td>7</td>
<td>TIM3/CD294</td>
<td>3</td>
</tr>
<tr>
<td>G-CSF</td>
<td>7</td>
<td>DR/11c (DC1)</td>
<td>3</td>
</tr>
<tr>
<td>IL-9</td>
<td>6</td>
<td>DR/123 (DC2)</td>
<td>3</td>
</tr>
<tr>
<td>VEGF</td>
<td>6</td>
<td>B7-H1(DRhi)</td>
<td>3</td>
</tr>
<tr>
<td>CD206</td>
<td>6</td>
<td>IL-7</td>
<td>2</td>
</tr>
<tr>
<td>IL-1ra</td>
<td>5</td>
<td>DR (hi)</td>
<td>3</td>
</tr>
<tr>
<td>IL-13</td>
<td>5</td>
<td>FGF</td>
<td>2</td>
</tr>
<tr>
<td>CD4/294</td>
<td>5</td>
<td>IP-10</td>
<td>2</td>
</tr>
<tr>
<td>CD11c/14</td>
<td>5</td>
<td>CD3/4</td>
<td>2</td>
</tr>
<tr>
<td>CD197/CD206</td>
<td>5</td>
<td>CD3/8</td>
<td>2</td>
</tr>
<tr>
<td>DR (hi)</td>
<td>5</td>
<td>CD4/TIM3</td>
<td>2</td>
</tr>
<tr>
<td>IL-17</td>
<td>5</td>
<td>B7-H1(DRlo)</td>
<td>2</td>
</tr>
<tr>
<td>IL-6</td>
<td>4</td>
<td>CRP pmol/L</td>
<td>1</td>
</tr>
<tr>
<td>IL-8</td>
<td>4</td>
<td>IL-1b</td>
<td>1</td>
</tr>
<tr>
<td>CD11c/CD123</td>
<td>4</td>
<td>TNF-a</td>
<td>1</td>
</tr>
<tr>
<td>Treg (% gated)</td>
<td>4</td>
<td>CD3/62L</td>
<td>1</td>
</tr>
<tr>
<td>IL-4</td>
<td>4</td>
<td>CD197</td>
<td>1</td>
</tr>
<tr>
<td>IL-5</td>
<td>3</td>
<td>MCP-1</td>
<td>0</td>
</tr>
<tr>
<td>GM-CSF</td>
<td>3</td>
<td>PDGF</td>
<td>0</td>
</tr>
<tr>
<td>MIP-1a</td>
<td>3</td>
<td>CD3/197</td>
<td>0</td>
</tr>
<tr>
<td>MIP-1b</td>
<td>3</td>
<td>DR (lo)</td>
<td>0</td>
</tr>
<tr>
<td>CD3/16+56</td>
<td>3</td>
<td>CD3/69</td>
<td>-1</td>
</tr>
</tbody>
</table>

was then calculated for each variable per individual patient. Patients were ranked by the sum of indices and the rank list was compared with the clinical outcome for each patient. Data suggest that patients in whom the fluctuation of measured immune parameters followed a well organized rhythm experienced the best clinical outcomes (PFS of 916 and 841 days for ranks 29 and 28, respectively) while the subjects with the lowest (-5 and -9) ranks (entirely random fluctuation of all measured immune variables) identified by this method had a PFS of 71 and 74 days, respectively (Figure 3). The latter were the only two patients in our cohort who had metastatic ocular melanoma. These patients not studied further given the disordered pattern of fluctuation and inability to fit them to any mathematical model.

Our analysis showed that CRP (empirically selected biomarker of immune oscillation) fit a time-dependent profile in less than 50% of the patients, and we found no correlation between PFS and CRP concentration and the first derivative (FD) of the fitted function on the day of treatment (Figure 4). Consequently, we selected for further study only the immune variables which fit a function in more than 50% of patients. In order to do this, we assigned a score to each immune variable which was obtained by summing up the indices across the remaining 8 patients with metastatic cutaneous melanoma (Table 2). Therefore, as the maximum theoretical score of a variable was 8 (8 patients), the score of 5 was chosen as cut-off because it selects variables which fit a function in more than half the patients. These included IL-1alpha, IL-9, IL-10, IL-12p70, IL-13, IL-15, IL-17, G-CSF, VEGF, Th2 T-helper lymphocytes (CD4/294), CD11c+ monocytes (CD11c/14), the ratio of polarized M1/M2 macrophages (CD197/CD206), and DR (hi) (cells that highly express HLA-DR). Since a large proportion of time dependent profiles were fitted to cosine curves using a somewhat non-stringent criterion (correlation coefficient), we further selected only data which fitted cosine curves with an R^2 > 75th percentile. The percentile value was calculated from the distribution of R^2 for curve fitting of all experimental profiles to the 9 mathematical functions across all patients. As a result, seven profiles for which the cosine function period was longer than the observation time (14 days) were eliminated. This yielded distinct time-dependent rhythms for the following immune parameters: the ratio of polarized M1/M2 macrophages (CD197/CD206), CD4/294, IL-12p70, IL-17, and CD11c+ monocytes (Figure 5). For multiple variables in 5 out of 8 patients these rhythms followed a predictable pattern which was a multiple of 3 days (3, 6, 9 and 12 days, respectively). A 4-day period was observed in one patient each for IL-12p70, IL-1ra and CD4/294. To better understand how the concentration or cell count and the trend for increase or decrease of these variables (FD of the fitted function) relate to clinical outcome, we compared the value of these variables in patients with different PFS. A fitted cosine curve was computed with all four parameters of the cosine function (a, b, c, and d: amplitude, period, phase shift and vertical shift) representing average values of the corresponding parameter across patients being compared and a variable being analyzed. The resulting curve represented averaged concentrations/cell count dynamics for several patients on a relative concentration scale (described above). First derivatives of the fitted function on the treatment day were also plotted on a relative scale for patients with different PFS. Concentration/cell counts and FD plots were constructed for CRP (Figure 4B), IL-12p70 and CD197/CD206 ratio (Figure 6 B, D) for those patients in whom these variables fitted a cosine function. The figures demonstrate that clinical outcome (PFS) is related to the concentration/cell count or the value of the FD of the given measurements (Figure 6).
Figure 3. Patient ranking based on ordered versus disordered pattern of fluctuation of serially measured immune parameters as related to clinical outcome (PFS). The ranks represent the sum of indices across 51 serially measured cytokine concentrations and immune cell counts for each patient over time; the maximum possible index value is 51.

Figure 4. (A) The extrapolated relative CRP concentration (right axis, dashed bars) and relative first derivative (FD) of the fitted function on the day of treatment (left axis, black bars) as related to PFS. “NF” marks cases where the time-dependent pattern of the variable dynamics did not fit any of the 9 mathematical functions. (B) Synthetic virtual curve of the averaged concentration dynamics of CRP for patients in whom data points fitted a cosine function. The curve represents a function where each of the four function parameters (amplitude, period, phase shift and vertical shift) were averaged for one variable across patients. Bars represent the relative value of the first derivative of the fitted function on the day of treatment. The numbers in callouts represent the PFS for each corresponding patient.
4.3. Identification of immunologically favorable time point in the immune response cycle (rhythm) for chemotherapy administration based on fluctuating systemic immune biomarkers

In terms of clinical application, our goal was to define a parameter which would encompass both the magnitude of change and the trend of fluctuation of a given variable at any time point in the immune response cycle, therefore describing time-dependent fluctuation of immune biomarkers more accurately than the protein concentration or cell count alone. For that purpose, we defined a new parameter (S) that was calculated as the sum of the relative concentration and FD with the formula: \( S = D + C \) where D is the relative first derivative and C is the relative concentration. In order to find the variables with the highest correlation between S and PFS on the day of treatment (with TMZ), we ranked the parameter S in descending order for each measured variable and estimated the association between the rank of each of the 14 variables which fit a function in more than half the patients and PFS. If a variable did not fit a biologically possible function, then the product could not be calculated (e.g.: infinity); since we analyzed 14 variables and the lowest rank for a product was 14, the next lowest rank for a product which could not be calculated was 15. Because this rank is weighted by the proportion of non-fitted variables in a given patient, we used a weighted rank calculated as 15*(number of variables which do not fit a function)/(total number of measured variables). Two variables, the concentration of IL12p70 and the ratio of CD197/CD206 positive cells (ratio of polarized M1/M2 macrophages) had the best association with PFS, which was further supported by estimation of association between the sum of ranks for these two variables and PFS (Figure 4). Fifty percent of patients (4/8) with the sum of ranks of IL12p70 and M1/M2 below 15 had an average PFS of 466 days, whereas the other 50% with a sum of ranks above 15 had an average PFS of 68 days, suggesting that the value of the parameter S on the day of treatment correlated favorably with clinical outcome in this limited data set. The parameter S on the day of treatment was 5.5 for the patient with the highest PFS (916 days) and 2.5 for the patient with the lowest PFS (37 days) suggesting that treatment administration at a time point in the cycle when S is elevated results in superior clinical response to therapy.

4.4. Full data integration and correlation with clinical outcome (clustering analysis)

Our preliminary data suggested that application of treatment at a time point when the parameter S is elevated may be associated with an improved clinical outcome. This trend was derived from the time-dependent profiles of 13 variables across 8 patients. The observed periodical pattern was consistent across profiles and implied the existence of periodicity of systemic immunity in patients with metastatic melanoma. However, the relationship between clinical outcome and the specific time point in the oscillation of each specific variable when treatment was applied varied. We therefore used a modified K-means clustering algorithm in order to develop more robust integral criteria for assessing the state of the immune system at a specific time point in the immune response cycle. With this method, the number of groups is determined from the number of full function periods which fit into one observation period. The algorithm computes the number of clusters for the entire range of integers from maximum to minimum numbers and outputs time points when the parameter S has the maximum value for each iteration (number of clusters) and for each variable. These time points are used as centroids for K- means clustering. Since the clustering results depend on the order of initial centroids, our modification performs clustering for all possible combinations of centroids and computes the date when the sum of indices for all clustered cosine profiles is maximal. Next, the algorithm computes the dates with maximum sums of relative S values across all possible combinations of centroids and numbers of clusters. These dates are outputted as optimal dates for chemotherapy application for a given patient and a given set of immune variables.

The algorithm divides the range of progression-free survival times into a number of bins ten times less than the number of patients. For each bin the algorithm counts the profiles of variables with R² above the cut-off value, and calculates the sum of S values on treatment start date for these variables. Next, linear regression analysis is performed both on the counts and on the sums, and the slope of the regression line is computed. Variables with high positive value of the sum of slopes have a positive correlation (PC) with PFS, variables with high negative value, a negative correlation (NEC), and variables with slopes close to zero have no correlation (NOC) with PFS. The cut-off for PC variables is the 75th percentile (mean + 0.67 x Standard Deviation) of all sum values and for the NEC the cut-off is the 25th percentile (mean - 0.67 x Standard Deviation). In this study we considered only the positive correlates.

In order to validate this approach, we next applied modified K-means cluster analysis to our clinical data and compared the optimal treatment time identified by this method with the actual timing of therapy delivery and the clinical outcome. The time profiles for IL12p70, IL-1ra, and CD206 satisfied the selection criteria from our pilot clinical trial data and were used for clustering. However, as only profiles which fit cosine functions with a correlation coefficient greater than 0.86 were used, IL-1ra was eliminated from clustering for patients #1, 4 and 7, and IL12p70 profile was eliminated for patient #10. Since only 6-7 data points were obtained prior to treatment initiation, we extrapolated 3 to 4 additional time points to match the same number of points (10) analyzed in the simulated data set (see below). The extrapolated values were computed using Fourier analysis. Clustering produced 1-3 days with maximum cumulative values of the parameter S for each patient (Figure 8). The optimal time for chemotherapy application predicted by the clustering algorithm fell within the 5-day period of actual chemotherapy delivery in the two patients with the longest PFS; in all other patients except one (patient # 12, in whom chemotherapy was started 4 days before the date that would have been predicted by this method), chemotherapy was applied several days before or after the optimal days predicted by our clustering algorithm (Figure 9).
4.5. The simulation model

In clinical practice it is often difficult to collect blood or tissue samples for long periods of time or frequently enough to obtain a time series of data points which unambiguously satisfy stringent curve fitting criteria and allow for mathematical modeling of clinical data. Therefore, given the maximum possible number of data points, we needed to determine the sampling frequency, observation period, curve amplitude and period (for periodical function) which fit a function with high probability and not only by chance alone. Consequently, we simulated time series of data points with input parameters derived from our pilot data. We considered 3 different observation periods (10, 15 and 20 days), 3 sampling frequencies (every day, every other day and 1-2 days), 100 amplitudes, and 20 periods. The variables in our study that fitted cosine curves by our selection criteria and had periods equal or shorter than 12 days were CD197/CD206 and IL12p70 (5 patients each), CD4/294 and IL-15 (4 patients each), CRP, IL-10, CD11c/14, CD206, IL-17, IL-13 (3 patients each); IL-1ra, IL-9, G-CSF and VEGF (2 patients each) and DR(hi) (one patient). Taking this into account, the amplitudes for a given variable were simulated as follows: the parameter B average (which defines the amplitude of the cosine function) was calculated across all patients in whom the time series for a particular variable fitted a cosine curve and the interval B_avg +/- two standard deviations was calculated and divided into 100 bins (Figure 10). Each of the 100 bin average was used in the cosine equation to produce profiles with particular amplitudes. Twenty different periods were simulated using the same technique; each data series was simulated with and without experimental error which was calculated using the coefficient of variation and maintaining the same distribution of error values as obtained in our clinical experiment. The error was added to or subtracted from the simulated values in random order. We simulated time series for the 16 variables which fitted cosine curves with R^2 above the 80 percentile cut-off in at least 7 out of 8 patients. Two sets of time series were simulated according
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Figure 6. (A, C) The extrapolated relative concentration (right axis, dashed bars) and relative first derivative (FD) of the fitted function on the day of treatment (left axis, black bars) as related to PFS. (A) IL-12p70; (C) CD197/CD206 ratio. “NF” marks cases where the time-dependent pattern of the variable dynamics did not fit any of the 9 mathematical functions. (B, D) Synthetic virtual curves of averaged concentration dynamics of IL12p70 (B) and CD197/CD206 (D) for patients in whom data points fitted a cosine function. The curve represents a function where each of the four function parameters (amplitude, period, phase shift and vertical shift) were averaged for one variable across patients. Bars represent the relative value of the first derivative of the fitted function on the day of treatment. Numbers in callouts represent the PFS for each corresponding patient.

to the described design. In the first set (Cosine profiles), the concentration/cell count values were calculated by the cosine formula. In the second set (Random profiles) values were produced by generation of random numbers within the set amplitude range. As a result, we obtained 576,000 data series of cosine profiles and 576,000 data series of random profiles, which were then fitted to the following five mathematical functions: logistic, quadratic, cosine, rational, Gaussian, and MMF function (Morgan-Mercer-Flodin) with R² recorded for each fitting. Next, we computed and analyzed the distribution of R² of the curve fitting in random and cosine data sets (Figure 11A). The analysis of the R² distribution for cosine function allows identification of the conditions (period, amplitude, sampling frequency, observation period, etc.) which predominantly produce true positive and true negative solutions, as well as those that produce false positive and false negative solutions. Of note, a solution is the conclusion whether or not the time series of data points fits a cosine curve based on the value of R². Simulated profiles computed by the cosine formula produced true positive and false negative solutions when R² was high or low correspondingly. Likewise, random profiles produced false positive and true negative solutions. Therefore, the ranges of R² values corresponding to high sensitivity and specificity of the solutions can be determined. This was necessary as one of the goals of our simulation study was to determine the cutoff values of R² which would allow us to achieve the best combination of specificity and sensitivity. A small number of time series (10185 profiles= 0.0088% of the total number of profiles) formed straight lines and were excluded from further analysis. For the cosine profiles, 81.7% (461998 out of 565821) of R² values were in the range 0.980 – 1.0; 50% of these profiles were obtained by fitting data series without introducing error and in the other fifty percent a simulated value of technical error was introduced. The 90th percentile of the R² values for the cosine profiles was 1.0 and 0.905 for the random profiles. The overall 90th percentile of the R² values in the range from 0 to 0.98 was 0.87. We then considered R² values in the range of 0.87 to 1.0. (Figure 11B) and accepted the 90th percentile of R² subset as the cut-off criteria for discriminating between a random set of data points and those calculated by the cosine formula. This cutoff (rather than the more stringent 0.98 cutoff) prevents obtaining a high number of false negative results. The resulting subset of R² values contains ambiguous solutions (false positive and false negative), the majority of which are introduced by profiles generated with observation periods of 10 days and every other day blood sampling frequency. When all profiles generated with both of these conditions were removed, only simulated cosine profiles fit cosine function with R² in the interval 0.870 to 0.995 (Figure 11C). No other tested observation period or sampling frequency produced a significant number of R² in this interval from random profiles. As expected, the proportion of R² above the 90th percentile cut-off obtained from fitting cosine profiles is higher for the profiles that have a higher number of time points, obtained through
longer observation periods or more frequent sampling, which is often a limiting factor in clinical practice. As a result, we used simulation studies to define the best clinical trial design that meets both the demands of curve fitting methods and clinical constraints. A schedule satisfying these conditions is 5 sequential days of blood sample collection followed by 2 days of rest and another 5 days of collection, which gives 6 degrees of freedom for data fitting to a cosine function. Further in the text we will refer to this schedule as 5-2-5. We next simulated time series for this schedule. All $R^2$ values (56,119 out of 56,128) above 0.980 were generated by fitting simulated cosine profiles (Figure 12 A, C). The $R^2$ obtained from fitting random profiles to cosine functions were largely in the range 0.000 to 0.980. The distribution of $R^2$ in this range is quasi-normal (Figure 12B) and the 90th percentile of this subset of $R^2$ values is 0.8055. Therefore, if 90th percentile is selected as a cut-off criteria for discriminating between the random set of data points and those calculated by the cosine formula, the ambiguous solutions will lie in the $R^2$ value range from 0.8055 to 0.980 (Figure 12D). When we performed the receiver operating characteristic (ROC) analysis of 16 variables for this interval of $R^2$ values, the best performing variable was IL-1ra (area under the curve (AUC) = 0.955) and the worst performing variable was CRP (AUC= 0.734).
Figure 9. Clustering of time-dependent profiles for IL-12p70 (red) and IL-1ra (black). The parameter S was calculated for each variable and clustering was performed by modified k-means clustering (see section 4.4) to find the optimal time for chemotherapy delivery on the days when the sum of the parameter S for the two variables was highest (vertical black lines). The day when chemotherapy was actually applied is represented by a yellow line. The analysis for patient # 1 (PFS= 916 days) is shown.

Figure 10. Data simulation design. 100 amplitudes (A) and 20 periods (B) were derived from the average values and standard deviations (SD) of measurements of these parameters in 5-7 patients for each of 16 variables (F). Data sets were simulated for cosine and for random distribution of data points. The simulation was performed for three different sampling frequencies (C), three observation periods (D), and with or without technical error of the measurement (E).
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Figure 11. The frequency distribution (log base 10) of R² obtained in the simulation study. The red bars represent frequencies for simulated cosine profiles; blue bars represent frequencies for simulated random profiles. (A) – Overall distribution of R² for cosine and random profiles. (B) - Frequency distribution of R² in the range between 0.87 (90th percentile) and 1. (C) - Frequency distribution of R² in the range between 0.87 and 1 excluding profiles generated for observation period of 10 days and every other day sampling frequency.

5. DISCUSSION

The presence of “biorhythms” in medicine has been a topic of extensive study for many years (23-25). The discovery that hormonal secretion is an episodic process opened a new era in endocrinology and revolutionized the way we now measure hormones. Rhythmic phenomena are typical for all levels of biological organization, with periods ranging from centuries and decades (evolution of species and ecological systems), to milliseconds of electrical potential in nerve and cardiac cells (26). Similar episodic changes have been described in immunobiology with patterns that follow a circadian (24 hour) (22), infradian (greater than 24 hours) (27), seasonal, or circannual (yearly) rhythm (23). Certainly, the pronounced circadian variation in T cell subpopulations in human blood during the 24-hour sleep-wake cycle is now well-documented (22,28) and seems to be controlled by the release of cortisol and catecholamines, the body’s major endocrinologic stress-hormone systems. Likewise, peripheral blood immune effectors (T cells and natural killer cells) fluctuate during normal menstrual cycle in healthy women, which may be critical for embryonic implantation and pregnancy (27). Our study describes similar time-dependent (dynamic) periodical fluctuations of systemic immunity in response to cancer in patients with metastatic melanoma. Our data also suggests that these immune biorhythms may be therapeutically relevant with respect to timing of chemotherapy administration and can have a dramatic impact on therapeutic outcome.

The idea of biorhythm-based delivery of therapy (chronopharmacology) has been a topic of interest for oncologists for many years (29,30). Chronotherapy implies that, by administering chemotherapy at the “right time of day”, one can take advantage of asynchronies in cell proliferation between malignant and normal cells, thereby minimizing chemotherapy toxicity and increasing treatment efficacy. However, despite the fact that most trials showed a difference in tolerability in favor of chronotherapy, so far the two largest randomized trials (31,32) showed no benefit in treatment efficacy. In the current study we extrapolated the principle of chronotherapy to utilize spontaneously developed immune infradian rhythms and this resulted in improved clinical response to chemotherapy, as predicted in the mouse models (18). The potential explanation is that application of chemotherapy at an immunologically favorable phase may selectively ablate the cycling
Figure 12. The frequency distribution of $R^2$ obtained in the simulation study for the 5-2-5 schedule. Red bars represent frequencies for simulated cosine profiles; blue bars represent frequencies for simulated random profiles. (A) - Overall distribution of $R^2$ for cosine and random profiles shows that all $R^2$ above 0.98 were obtained from simulated cosine profiles. (B) - Frequency distribution of $R^2$ in the range between 0 and 0.99 shows normal distribution of $R^2$ obtained from random profiles. (C) - Frequency distribution of $R^2$ in the range between 0.98 and 1 demonstrates that the number of $R^2$ obtained from random profiles is insignificant in this range. (D) - Frequency distribution of $R^2$ in the range between 0.8055 (90th percentile) and 0.98 demonstrates that ambiguous solutions (false positive and false negative) are frequent in this range.

Suppressive elements of immunity and release the patients’ immune system from down-regulation. The fluctuation of CRP concentration was an attractive candidate for timing of chemotherapy given its well established quantification methodology (33,34) and previously described periodic fluctuations in healthy individuals (35), patients with chronic viral infections (36) or cancer (10). However, our data suggested that this approach may be overly simplistic, as it did not take into account the complex and dynamic interaction among the multiple fluctuating biomarkers that define the systemic anti-tumor immune response. The accurate prediction of the optimal time for chemotherapy administration likely requires a global analysis of the changes of systemic immunity over time for each individual patient. Using mathematical modeling and curve fitting analysis we defined the parameter ($S$) that encompasses both the magnitude of change in concentration or cell count and the trend for increase or decrease of a given immune biomarker at a particular time-point in the anti-tumor immune response cycle. Our preliminary data suggested that application of treatment at a time when maximum numbers of immune variables have maximum cumulative value of the parameter $S$ could result in improved clinical outcomes. This hypothesis was corroborated by the application of modified K-means clustering analysis to our clinical trial data. We found that
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the time of chemotherapy application predicted by the clustering algorithm fell within the 5-day period of actual chemotherapy application only in the two patients with the longest PFS, while virtually all other patients received chemotherapy several days before or after the optimal days predicted by clustering. The two patients (25%) that were accurately selected for timing of therapy by our method had a complete response (CR) to treatment delivered based on the studied immune biorhythms. This response rate is much higher than the CR rate of 3% of TMZ treated historical controls (37).

It has been known for some time that disruption of biological rhythms may have clinical consequences. For instance, carefully designed observational studies have shown that people whose circadian rhythms are chronically disrupted are more prone to developing cancer (e.g. night-shift workers and breast cancer risk) (38, 39). Likewise, cancer growth may disturb biorhythms in the host (40, 41). These observations are consistent with our findings that patients with a disorganized (non-curve-fitting) anti-tumor immune response experienced a decreased disease-free survival (average PFS of 69 days), relative to those in whom the measured immune parameters followed a predictable biorhythm (average PFS of 465 days).

Surprisingly, the subjects with an entirely random fluctuation of cytokine concentrations/cell counts identified by this method were the only two patients with metastatic ocular melanoma enrolled in this study. Ocular melanoma has previously been shown to differ markedly from other melanoma subtypes both in clinical behavior and molecular makeup and our findings raise the question whether these patients may have a distinctive immunological signature as well. On the other hand, the best clinical responses were observed in the two patients who maintained a well synchronized anti-tumor immune response, possibly overcoming the global immune dysfunction of malignancy. Timed delivery of chemotherapy in that context may have allowed for a more precise therapeutic intervention leading to putative depletion of immune down-regulatory signals in favor of effective anti-tumor immunity.

While cytotoxic anticancer drugs are intended to primarily affect cancer cells directly, many such agents also exhibit "side effects" impacting cells of the immune system (bone marrow). Hematologic (bone marrow) toxicity of cytotoxic chemotherapy is the most common side effect of modern anti-cancer therapy and the principal determinant of the design of cancer chemotherapy treatment schedules. This common (nearly universal) side effect of chemotherapy is seldom considered in the interpretation of clinical efficacy for a given anti-cancer agent. Although the underlying rationale is not yet certain, it is possible that the common hematologic toxicities of TMZ yield alterations of systemic immunity which may influence clinical outcome, as has been previously suggested (42). This effect appears to be related to Treg function (17, 42). Pharmacokinetic studies have demonstrated that PBMC exposed to five subsequent TMZ treatments (concentration and schedule comparable to that obtainable in the clinic) exhibited a significant inhibition of their proliferative response (43). Although TMZ administration would theoretically inhibit desirable effector T cells as well, we propose that timed administration of this agent may selectively suppress Treg who lag behind T effectors in their clonotypic expansion (17). By that time, effector T cells would have proliferated and become activated and may be therefore less susceptible to the effects of TMZ chemotherapy. Clearly, administration of the same agent (TMZ) at the same dose/schedule only delivered at a different time in the "immune biorhythm" of the patient may have an entirely different cumulative effect (depletion of clonotypic tumor specific CTL) on immune homeostasis and possibly on clinical outcome.

Albeit intriguing and suggestive, our present data can only be interpreted as hypothesis generating due primarily to the small number of patients and specimens studied. At the time of the original design of this study we had no appreciation of the underlining immune biorhythms and therefore had to design a clinical trial in which we empirically selected an arbitrary pre-treatment observation period (14 days) in which we would collect an arbitrary number of blood samples (5-8) collected at arbitrary intervals (every 2-3 days). The presented data have now given us the necessary information to design a follow-up clinical trial that will be better suited to truly test the hypothesis proposed herein. The development of a strong mathematical model, central to our understanding of dynamic processes requires an adequate number of serial measurements. Fitting of 6 or 7 data points to a function with four parameters (sinusoidal and rational functions) can be ambiguous even if the goodness-of-fit metrics are satisfactory ($R^2$ and coefficient of variation are close to 1.0, confidence interval is narrow.). Therefore, in order to design a sample collection schedule which permits obtaining a sufficient number of time points but keeps the blood collection burden as low as possible for our patients, we used simulation and modeling studies for different observation periods and sampling frequencies. The goal was to identify a clinical trial design which meets both the analytical (mathematical modeling) and clinical (burden to patient) requirements. The result of these analyses suggested that a schedule satisfying these conditions is 5 sequential days of blood sample collection followed by 2 days of rest and another 5 days of collection (5-2-5 schedule) yielding sufficient statistical power to address the question at hand. Based on these findings we were able to design a clinical trial that directly tests the hypothesis generated by our presented data which suggests that synchronizing conventional chemotherapy delivery with pre-existing, patient-specific (individualized) biorhythms, may yield improved clinical outcomes using mechanisms believed to only be a toxic side-effect of therapy (lymphodepletion). Said validation study is currently under way.

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**Abbreviations:** CRP: C-reactive protein ; Treg: T-regulatory cells; Th cells: T-helper cells; TMZ: temozolomide; IL: interleukin; GFG: basic fibroblast growth factor (FGF); G-CSF: granulocyte colony-stimulating factor; GM-CSF: granulocyte-macrophage colony-stimulating factor; IFN-gamma: interferon gamma; IP-10:interferon-gamma-induced protein; MCP-1: macrophage chemoattractant protein 1; MIP-1alpha: migration inhibitory protein 1alpha; PDGF: platelet-derived growth factor; RANTES: Regulated upon Activation Normal T-cell Expressed and Secreted; TNF-alpha: tumor necrosis factor alpha; VEGF: vascular endothelial growth factor ; TGF-beta1: Transforming growth factor beta; PBMC: peripheral blood mononuclear cell; RECIST: Response Evaluation Criteria in Solid Tumors; ECOG: Eastern Cooperative Oncology Group;
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FD: first derivative; CC: correlation coefficient; PFS: progression-free survival

Key Words: Cancer, Biorhythm, Infradian, Inflammation, Oscillation, Dynamics, Cytokines, Immune Cells, Computer Simulation

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