Digital Display Precision Predictor: the prototype of a global biomarker model to guide treatments with targeted therapy and predict progression-free survival

Vladimir Lazar1,8, Shai Magidi1,18, Nicolas Girard2, Alexia Savignoni2, Jean-François Martin1, Giorgio Massimini4, Catherine Bresson1, Raanan Berger5, Amir Onn5, Jacques Raynaud6, Fanny Wunder1, Ioana Berindan-Neagoe7,8, Marina Sekacheva9, Irene Braha10, Josep Tabernero10, Enriquea Felip10, Angel Porgador11, Claudia Kleinman12, Gerald Batist12, Benjamin Solomon13, Apostolia Maria Tsimberidou14, Jean-Charles Soria15, Eitan Rubin1,18, Razelle Kurzrock16,18 and Richard L. Schilsky17,18

The expanding targeted therapy landscape requires combinatorial biomarkers for patient stratification and treatment selection. This requires simultaneous exploration of multiple genes of relevant networks to account for the complexity of mechanisms that govern drug sensitivity and predict clinical outcomes. We present the algorithm, Digital Display Precision Predictor (DDPP), aiming to identify transcriptomic predictors of treatment outcome. For example, 17 and 13 key genes were derived from the literature by their association with MTOR and angiogenesis pathways, respectively, and their expression in tumor versus normal tissues was associated with the progression-free survival (PFS) of patients treated with everolimus or axitinib (respectively) using DDPP. A specific eight-gene set best correlated with PFS in six patients treated with everolimus: AKT2, TSC1, FKB-12, TSC2, RPTOR, RHEB, PIK3CA, and PIK3CB (r = 0.99, p = 5.67E−05). A two-gene set best correlated with PFS in five patients treated with axitinib: KIT and KITLG (r = 0.99, p = 4.68E−04). Leave-one-out experiments demonstrated significant concordance between observed and DDPP-predicted PFS (r = 0.9, p = 0.015) for patients treated with everolimus. Notwithstanding the small cohort and pending further prospective validation, the prototype of DDPP offers the potential to transform patients’ treatment selection with a tumor- and treatment-agnostic predictor of outcomes (duration of PFS).

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INTRODUCTION

The application of personalized medicine to oncology has resulted in prominent successes that have led to approved, molecularly specific, biomarker-defined indications for targeted therapies. For example, the use of EGFR mutation/erlotinib7, KIT mutation/imatinib2, BRAF mutation/ vemurafenib3, ALK translocation/crizotinib4, and high tumor mutation burden or microsatellite instability high/pembrolizumab5-8 have dramatically changed the treatment landscape in many cancers including, but not limited to melanoma, non-small cell lung carcinoma (NSCLC), colorectal carcinoma (CRC), and head and neck (HN) cancers.

However, despite the advent of personalized precision oncology, cancer remains one of the leading causes of deaths all over the world. Globally, 9.6 million deaths are attributed to cancer, representing 13% of all deaths5.

Extending the application of precision medicine requires a deeper understanding of tumor biology. Furthermore, improvement in the ability to select patients is needed, both with respect to identifying sensitive versus resistant tumors and in pinpointing each patient’s tumor for all markers related to all possible targeted therapies becomes infeasible due to the limited amount of tissue usually obtained by biopsies, pointing out the limitation of the classic companion diagnostic approach.

A comprehensive analysis of all relevant genes in a single assay would enable the exploration of all drug targets simultaneously to inform therapeutic options for patients. Furthermore, the complexity of cancer biology requires investigation of multiple genes in networks of pathways to understand the variability of clinical outcomes observed, that cannot be achieved by investigating single genes (as performed with most companion diagnostic tests).

We report the Digital Display Precision Predictor (DDPP), a biomarker strategy and tool, able to predict duration of progression-free survival (PFS) for multiple targeted treatments, based on the comprehensive investigation of the whole transcriptome. The DDPP prototype presented here was derived from analysis of transcriptomic and clinical outcomes in patients with advanced cancer enrolled in the WINCHER clinical trial and treated with: everolimus (mTOR inhibitor), axitinib (VEGFR receptors inhibitor), afatinib (pan-HER inhibitor), trametinib (MEK inhibitor), FGFR inhibitors, and anti-PD-1/PD-L1 antibodies (pembrolizumab, nivolumab and atezolizumab). The WINCHER trial (NCT...
Indeed, the two patients with the longest PFS, both with GI tract neuroendocrine tumors, had no mutations (ID 203) or no genomic alterations in the PI3K/AKT/mTOR pathway (ID 148), respectively. In contrast, the patients with much shorter PFS (ID 227, ID 6, ID 90, and ID 117) did have alterations in the PI3K/AKT/mTOR pathway, albeit accompanied by co-alterations that might have driven resistance. Since DNA biomarkers could not explain variations in clinical outcome, we further investigated whether transcriptomics and DDPP could provide a deeper insight. Figure 1 shows the DDPP profile of the 17 key genes of the six patients treated with everolimus as monotherapy. It should be noted that the DDPP profile is highly variable between the six patients. While MTOR has the same steady-state level in both tumor and normal tissues, Fig. 1 e, f suggests a trend of greater overexpression of upstream regulators of MTORC1 complex (in particular TSC1 and AKT2) in patients with longer PFS (ID 148 and ID 203), as compared with the patients with shorter PFS (ID 090, ID 227, ID 117, and ID 006).

The relative impact on the PFS of each individual gene was determined through Pearson correlations between differential gene expression and PFS for each of the 17 genes were AKT2 (r = 0.75, p = 0.087), TSC1 (r = 0.74, p = 0.094), FKB-12 (r = 0.67, p = 0.149), TSC2 (r = 0.63, p = 0.178), RPTOR (r = 0.61, p = 0.198), RHEB (r = 0.49, p = 0.325), PIK3CA (r = 0.43, p = 0.4), PIK3CB (r = 0.41, p = 0.414), AKT1 (r = 0.35, p = 0.496), MLST8 (r = 0.34, p = 0.509), VEGFA (r = 0.27, p = 0.604), HIF1 (r = 0.27, p = 0.606), PTEN (r = 0.16, p = 0.759), 4EBP1 (r = 0.16, p = 0.77), RICTOR (r = 0.14, p = 0.788), MTOR (r = 0.13, p = 0.807), and S6K1 (r = 0.04, p = 0.936). We further explored the combined differential expression in tumor versus normal tissues of the most contributive key genes involved in the everolimus pathway. For each of the correlations with PFS, we built a vectorial summation using a “step-in” method, starting with AKT2, which was identified as the most contributive gene according to the Pearson correlations, and adding successively a single gene at each step in the order of their significance: AKT2-TSC1; AKT2-TSC1-FKB12, then AKT2-TSC1-FKB12-TSC2 and so forth, obtaining in total 17 different vector summations. Each combined vector was correlated with the observed PFS.

Figure 2a shows that the optimal performance was obtained by the vectorial summation of the eight most contributive genes AKT2, TSC1, FKB-12, TSC2, RPTOR, RHEB, PIK3CA, and PIK3CB, which showed the most significant correlation with PFS, among the 17 possibilities: (r = 0.99, p = 5.67e−05). The higher the relative expression of these key genes in tumor tissues, the longer the PFS is under treatment with everolimus. The linear regression model for the correlation with PFS is $Y = 1.499e - 13X + 3.134$ (Equation 1), where $X =$ the absolute value of the fold of log2(fold change tumor versus normal) multiplied by log1.1 (Intensity_Tumor) of each value for each of the eight genes, and $Y =$ PFS in months. Allelic frequency of mutations was not taken into account, as mutations could not explain variations in PFS. Allelic frequency was not available as transcriptomics was performed with microarray technology. It should be noted that the most contributive genes, AKT2, TSC1, FKB-12, TSC2, RPTOR, and RHEB are key for direct interaction with MTOR and its upstream regulation. The linear regression model for the correlation with PFS is $Y = 1.499e - 13X + 3.134$ (Equation 1), where $X =$ the absolute value of the fold of log2(fold change tumor versus normal) multiplied by log1.1 (Intensity_Tumor) of each value for each of the eight genes, and $Y =$ PFS in months. Allelic frequency of mutations was not taken into account, as mutations could not explain variations in PFS. Allelic frequency was not available as transcriptomics was performed with microarray technology. It should be noted that the most contributive genes, AKT2, TSC1, FKB-12, TSC2, RPTOR, and RHEB are key for direct interaction with MTOR and its upstream regulation (TSC1, TSC2, and RHEB). Furthermore, FKB-12 binds everolimus and associates to MTOR forming together with RPTOR the MTORC1 complex.

Figure 2b shows that when only the tumor biopsy was investigated (as is usually done in the current biomarker studies in oncology practice or translational research), the significance of the correlation dropped to $r = 0.24$, $p = 0.65$, suggesting the critical importance of the new strategy of assessing tumor versus normal analogous organ-matched dual biopsy.

In order to assess the prognostic versus the predictive value of the DDPP data in our analyses, we tested the specific predictor of the PFS for everolimus (n = 6 patients) generated by eight genes (AKT2, TSC1, FKB-12, TSC2, RPTOR, RHEB, PIK3CA, and MTOR) was feasible and increased, by about one third, the number of patients that could be matched to a targeted therapy as compared to genomic analysis alone.

RESULTS

DDPP strategy and objectives

The objective of the DDPP was to build a predictor of the PFS as a continuous variable, in contrast with current binary predictive biomarkers that predict if patients will respond to a specific therapeutic regimen or not. The DDPP transcriptomic-based biomarker strategy is based on three pillars: (1) gene selection—literature review was used to identify a list of key genes governing the sensitivity for a given therapeutic regimen and the exploration of their differential expression in tumor compared to analogous normal tissues. Comprehensive transcriptomic analysis of the whole transcriptome (~20,000 genes) in a single assay requiring a very small amount of tissue, which enables the exploration of all targets simultaneously to inform therapy options for patients. Reducing the number of features (genes) for consideration reduces the chances of overfitting and allows useful predictions to be made regardless of the size of the patient cohort; (2) ranking each of the genes from the identified list of relevant genes, based on their relative association with PFS. Different conventional ranking methods were considered, including Cox univariate and multivariate regression models, multiple linear regression (MLR), or parametric (Pearson) and nonparametric (Spearman) correlation tests. We selected the Pearson correlation with PFS, since it outperformed all other ranking methods (presented in “Methods” and in Supplemental Notes); and (3) building a PFS predictor. Once the relevant genes were ranked, we applied an empirical summation approach, comparing the performance of every set of genes with the top K genes ($K = 1...N$, where $N$ is the number of genes in the list identified in step (1) above) in predicting PFS. For every subset with the top-ranking K genes, four summation methods were used to predict PFS: mean, median, product, and absolute product.

DDPP investigations of patients treated with everolimus

We applied the DDPP method to build a PFS predictor for everolimus treatment: Supplemental Table 1 describes the currently recognized 17 key genes of the mTOR pathway11,12: upstream regulators of MTOR: PIK3CA, PIK3CB, AKT1, AKT2, PTEN, TSC1, TSC2, RHEB, and FKB-12 (FKB1PA) plays a key role as it binds to everolimus and interacts with MTOR, resulting in the formation of the inhibitory complex MTORC1 (MTOR, MLST8, and RPTOR) and MTORC2 (MTOR, MLST8, and RICTOR); downstream effectors: S6K1 (RSK6KB1), 4EBP1 (EIF4EBP1), HIF1 (HIF1A), and VEGFA.

In six patients treated with everolimus, the fold changes measuring the differential tumor versus normal gene expression of each of the 17 key genes, creates 17 different coordinates defined by the fold changes in tumor and the steady-state levels of mRNAs in tumor and normal tissues. The main clinical and outcome characteristics of the patients treated with everolimus are described in Table 1, together with next-generation sequencing Foundation One test (Foundation Medicine) performed during the WINOTHER study. Results of the WINOTHER trial have been previously published16. One patient treated with everolimus (ID 203) had an exceptional response lasting in excess of 60 months and PFS was therefore censored at this timepoint. It should be noted that the genomic profile could not explain the variation in PFS observed in the everolimus monotherapy group;
and PIK3CB) and cross-correlated their combined differential expression with the PFS of patients under axitinib treatment (n = 5, Table 1). Figure 2c demonstrates that the subset of genes identified for everolimus was highly specific to the everolimus regimen, as the cross correlation with PFS of patients treated with axitinib was not significant (r = 0.29, p = 0.637), suggesting that the DDPP findings are consistent with the known biology of the MTOR pathway and the mechanism of action of everolimus.

To interrogate whether the findings of these analyses could be used as predictors, and address the potential for overfitting, we performed leave-one-out analyses, reiterating six combinatorial analyses. At each investigation, one patient was discarded, and a correlator/predictor was identified based on the remaining five patients applying the same methodology. The correlator was then used as a predictor to predict the PFS of the patient left out. Figure 2d depicts the six correlations obtained in the reiterative experiments, and Fig. 2e shows the correlation between the observed PFS and the leave-one-out predicted PFS. At each reiteration, the predictor is generated by different subsets of experiments, and Fig. 2e shows the correlation between the observed PFS and the leave-one-out predicted PFS. At each reiteration, the predictor is generated by different subsets of experiments, and Fig. 2e shows the correlation between the observed PFS and the leave-one-out predicted PFS.

Table 1. Characteristics of the patients treated in WINTHER trial investigated with DDPP.

| Study ID | Age | Sex | Cancer site | Prior lines | PFS months | DNA—list of molecular alterations (Foundation Medicine report)* | Drug_given |
|----------|-----|-----|-------------|-------------|------------|---------------------------------------------------------------|------------|
| 203      | 67  | F   | GI/NE       | 1           | 60.0+      | No mutation                                                  | Everolimus |
| 148      | 82  | M   | GI/NE       | 2           | 11.6       | BCOR N1652fs*34; CDKN1B E126fs*1                              | Everolimus |
| 6        | 64  | F   | UP          | 1           | 8.1        | TSC1 splice site 913 + 1G>T; BRCA1 truncation, intron 11; CDKN2A B loss; DNMT3A R882H; LRPR1 B loss | Everolimus |
| 117      | 34  | M   | HN          | 2           | 1.9        | TSC2 S1431L; TP53 G2455; BCOR K374fs*19; SMARCA4 R1135W       | Everolimus |
| 227      | 56  | M   | LS          | 4           | 8.8        | MTOR L2209V; ETV6 trunc intron 5; CIC S333fs*36; MLL2 G3698 fs*51 | Axitinib   |
| 83       | 59  | M   | HN          | 4           | 8.8        | MTOR L2209V; ETV6 trunc intron 5; CIC S333fs*36; MLL2 G3698 fs*51 | Axitinib   |
| 223      | 65  | F   | HN          | 3           | 7.1        | CCND1 T2861                                                  | Axitinib   |
| 259      | 53  | F   | HN          | 4           | 6.2        | PDGFRB amp                                                   | Axitinib   |
| 25       | 65  | M   | LN          | 2           | 5.3        | TP53 N1995F; KDM3A T275fs*4; MSH6 K1358fs*2; NFE2L2 R18Q     | Axitinib   |
| 88       | 56  | M   | Lung        | 1           | 2.9        | DNMT3A R635P; KRAS G12C; TP53 Y220C; MLL2 T1246M              | Axitinib   |
| 149      | 54  | F   | CRC         | 5           | 7.4        | KRAS G12V; ARID1A SPlice SITE 2733-1G>A                      | Trametinib |
| 100      | 43  | M   | CRC         | 2           | 6.6        | NAB2 F999insT; IDH1 R132C                                    | Trametinib |
| 118      | 78  | F   | Lung        | 3           | 3.1        | KRAS G12C; CDKN2A B loss; TP53 V157F; Y220 fs*27; MTHY1 G382D | Trametinib |
| 156      | 71  | F   | Lung        | 2           | 14.3       | EGFR E746; A750del; T790M; CDKNA2 B loss; CTNNB1 S333F; MYC amplification; SMAD4 P186fs*6; STAG2 splice site 1535-12_1630del108 | Afatinib + cetuximab |
| 235      | 60  | F   | Lung        | 1           | 11.3       | ERBB2 A775_G776insYVMA                                       | Afatinib   |
| 136      | 79  | M   | Lung        | 3           | 0.4        | ERBB3 amp; MET splice site 3028 + 1G>A; STK11 Q1000 ATM        | Afatinib   |
| 237      | 47  | M   | HN          | 6           | 19.3       | CCND1 amp; FGFR2 amp; CDKN2A B loss; FGFR1 amp; TP53 V157F; Y220 fs*27; MTHY1 G382D | NCT01004224 BGJ398 |
| 247      | 67  | M   | Esophagus   | 2           | 1.6        | FGFR2 amp; CDKN2A B loss; TP53 W91*; ASXL1 splice site 472-2A > G | NCT02052778 TAS-120 |
| 228      | 38  | M   | CRC         | 5           | 0.7        | FGFR1 amp; TP53 C176F; APC E1322*; R213*; SMAD4 loss; SOX9 V163fs*21 | NCT02052778 TAS-120 |
| 183      | 66  | M   | CRC         | 2           | 61.0+      | RBB3; V104M; MAP2K1; E203K; CDKN2A B loss; FBXW7 R465C; PIK3CA E39K; PIK3R1 R348*, R639*; PTEN R233*; splice site 801 + 2T > G; TP53 R158H; R273H; APC R1450*; R499*; ARID1A P1115fs*46, Q1306fs*17; ATRX Q2422; CDH1 D433N; SMAD4 P186fs*6; STAG2 splice site 1535-12_1630del108 | Pembrolizumab (TMB: 74.8) (MSI: +) |
| 294      | 57  | M   | HN          | 1           | 1.7        | BRCA2 K3408*                                                | Nivolumab (TMB: 0) (MSI: −) |
| 270      | 76  | F   | CRC         | 3           | 0.9        | FLT4 amp; FLT3 amp equivocal; BARD1 C53fs*5; MYC amp; PARK2 loss exons 3-5; TP53 R1755; APC T1556fs*3; BCL2L1 amp; CDK8 amp; ETV6 rearrangement intron 5; FAM123B R497*; GATA6 amp equivocall; KDM6A-Y215*; MTHY1 G382D; NOTCH1 Q2123* | Atezolizumab (TMB: 10.4) (MSI: −) |

Foundation Medicine®; ID 203, PFS 60+; and OS60+ are censored values, updated from the WINTHER trial Supplemental Table 3. Abbreviations: GI gastrointestinal, NE neuroendocrine, HN head and neck, UP unknown primary, LS liposarcoma, TMB tumor mutation burden, MSI microsatellite instability, amp amplification, del deletion, trunc truncation.
This observation pinpoints the need to validate the DDPP on a larger cohort of patients. Nevertheless, the concordance between the real PFS of the patients left out, and the predicted PFS using the correlator obtained at each reiteration on the remaining five patients is significant, with \( r = 0.9, p = 0.015 \).

We evaluated the performance of the predictive method by an independent method assessing the root mean squared error (RMSE) that was 16.82. The mean absolute error that measures the direction of the error unlike the RMSE was 9.32. Given the small magnitude of the error without taking into consideration the number of patients used as training and test datasets, and (ii) DDPP could potentially be used for identification of new genes/biomarkers, if used not in a random selection manner, but rather in a systematic selection manner.

**DDPP tested as a biomarker discovery method.** The most significant correlation between the differential expression of a new identified gene and PFS of patients treated with everolimus was observed for **UBAP1** (ubiquitin associated protein 1): \( r = 0.99, p = 7.56E−07 \) (Fig. 2f). **UBAP1** (NAG20) is a component of the ESCRT-I complex, a regulator of vesicular trafficking that could be potentially involved in degradation of MTORC2 and directly linked to MTOR pathway.\(^1\)

**DDP and prediction of outcome with axitinib**

At nanomolar concentrations, axitinib specifically inhibits VEGFR1, VEGFR2, and VEGFR3. Thirteen key genes involved in the control of angiogenesis were selected and investigated with DDPP methodology: **FLT1** (VEGFR1), **KDR** (VEGFR2), **FLT4** (VEGFR3), and their ligands **VEGFA**, **VEGFB**, **VEGFC**, and **FIGF**, **PDGFRB**, **PDGFA**, **PDGFB**, **KIT**, and **KITLG**.\(^15,16\) Four patients had **HN** carcinoma, and one patient had a lung adenocarcinoma. Table 1 shows the observation that randomly selected sets of genes generated significant correlations with PFS suggests that (i) specificity of correlations could be increased only with a larger number of patients used as training and test datasets, and (ii) DDPP could potentially be used for identification of new genes/biomarkers, if used not in a random selection manner, but rather in a systematic selection manner.
different PFS under treatment with axitinib observed in the
WINHER trial.

The differential tumor versus normal expression of KIT and of its ligand KITLG was identified as being the major driver of the correlation with the PFS of the patients treated with axitinib (Supplemental Fig. 1a, b), and their combined vector KIT-KITLG generated the optimal performance in the correlation with PFS: \( r = 0.99, p = 4.68 \times 10^{-4} \) (Supplemental Fig. 1c). The linear regression model is \( Y = 2.014 X - 0.02 X^2 + 4.36 \) (Equation 2), where \( X \) is the sum of log2(fold change tumor versus normal) multiplied by log1.1 (Intensity_Normal) of each value for each of the two genes, and \( Y = 0.1 \) is the PFS in months.

Random selections of two genes across the whole transcriptome and correlation of their vectorial summation with PFS of the five patients treated with axitinib was repeated 100,000 times. Using the same threshold as the specific predictor \( r = 0.99, p = 4.68 \times 10^{-4} \) the percentage of significant correlations was 0.059%.

Leave one out experiments. We performed (using the same “step-in” vectorial summation methodology) five leave-one-out reiterations, discarding at each experiment one patient and building a predictor on the remaining four to address potential overfitting. We observed again an instability of the predictors and dependence on the composition of the cohorts at each reiteration. The concordance between real PFS of the patients left out, and the predicted PFS using the correlator obtained at each reiteration was lower than for the everolimus example \( r = 0.81, p = 0.1 \) likely related to a lower number of patients in each reiteration. These data suggest again that performance and accuracy of the prediction of the PFS could be increased only with a higher number of patients in the training and validation datasets.

DDP and other examples of TKIs PFS predictors

Trametinib. Thirteen key genes were investigated: MEK1 (MAP2K1), MEK2 (MAP2K2), ARAF, BRAF, RAF1, ERK1 (MAPK3), ERK2 (MAPK1), MAPK10, KRAS, HRAS, NRAS, KSR1, and RAP1A. The combined differential tumor versus normal tissue expression of nine genes and their vectorial summation correlated with the PFS of three patients treated with trametinib as monotherapy (Table 1 and Supplemental Fig. 2a–c): ERK2, ARAF, CRAF, MEK1, MEK2, HRAS, ERK1, MAPK10, and KSR1 \( r = 0.99, p = 0.026 \); the linear regression model is \( Y = -6.872 X - 15X + 7.745 \) (Equation 3), where \( X \) is the fold of log2(fold change tumor versus normal) multiplied by log1.1 (Intensity_Tumor) for each of the nine genes, and \( Y = 0.1 \) is the PFS in months.

Afatinib. Thirteen key genes were investigated: EGFR, ERBB2, ERBB3, ERBB4 and their ligands EGF, TGFα, AR, EG, HBEFG, BTC, NRG1, NRG2, and NRG4. The combined differential tumor versus

Fig. 2 DDPP correlation with PFS for patients treated with everolimus. a Pearson correlation plots of the eight-gene predictor: AKT2, TSC1, FKB-12, TSC2, RPTOR, RHEB, PIK3CA, and PIK3CB with the PFS of six patients treated with everolimus as monotherapy; one patient out of these six had censored PFS (ID 203). X axis: absolute fold value of log2 based fold-changes tumor versus normal multiplied by log 1.1 based of the intensities in normal values for each of the eight genes selected; Y axis: PFS under treatment with everolimus as monotherapy in months. b Pearson correlation plots of the eight-gene predictor: AKT2, TSC1, FKB-12, TSC2, RPTOR, RHEB, PIK3CA, and PIK3CB with the PFS of six patients treated with everolimus as monotherapy when only the tumor biopsy is investigated; X axis: absolute fold value of log2 based of the intensities in tumor values for each of the eight genes selected; Y axis: PFS under treatment with everolimus as monotherapy in months. c Shuffle experiment: Pearson correlation plots of the 8 gene specific predictor for everolimus (AKT2, TSC1, FKB-12, TSC2, RPTOR, RHEB, PIK3CA, and PIK3CB) with the PFS of five patients treated with axitinib as monotherapy (Table 1); X axis: absolute fold value of log2 based fold-changes tumor versus normal multiplied by log 1.1 based of the intensities in normal values for each of the eight genes selected; Y axis: PFS under treatment with axitinib as monotherapy in months. d Leave-one-out experiments: each reiteration generates a predictor used to calculate PFS of the patient discarded. e Pearson correlation between leave-one-out predicted PFS and the observed PFS; X axis: predicted PFS as defined by leave-one-out (in months); Y axis: PFS under treatment with everolimus as monotherapy in months observed in WINHER trial. f Pearson correlation plot for UBA1 (NAG20) with the PFS of six patients treated with everolimus. X axis: log2 based fold-changes tumor versus normal multiplied by log 1.1 based of the intensities in tumor; Y axis: PFS under treatment with everolimus as monotherapy in months.
normal tissue expression of two genes and their vectorial summation correlated with the PFS of three patients treated with afatinib as monotherapy (Table 1 and Supplemental Fig. 2d–f). NRG4 and NRG2 (r = −1 p = 8.4E−04). The linear regression model is 

\[ Y = -4.558 - 0.360X + 2.549 \]  

(Equation 4), where \( X \) = the sum of log2(fold change tumor versus normal) multiplied by log1.1(Intensity_Tumor) of each value for each of the two genes, and \( Y \) = PFS in months.

**FGFR inhibitors.** Nineteen key genes investigated: FGFR1, FGFR2, FGFR3, FGFR4, and the FGFR ligands 1, 2, 3, 4, 5, etc.). The differential expression and vectorial summation of five genes correlated with the PFS of three patients treated with FGFR inhibitors BGJ398 or TAS-120 as monotherapy (Table 1 and Supplemental Fig. 2g–i): FGFR10, FGFR16, FGFR5, FGFR2, and FGFR13 (r = −1 p = 3.27E−03). The linear regression model is 

\[ Y = -5.273 - 0.20X + 5.135 \]  

(Equation 5), where \( X \) = the sum of log2(fold change tumor versus normal) multiplied by log1.1(Intensity_Normal) of each value for each of the five genes, and \( Y \) = PFS in months.

### DDPP and prediction of outcome after IO treatment

Although the most advanced knowledge has been generated around the therapies targeting PD-1/PDL-1 or CTLA-4, there are multiple other important pathways that may impact the immune response to cancer involving many genes, in particular LAG-3, TLR-4, VISTA, TIM-3, TIGIT, ICOS, OX40, and GITR. Among them LAG-3 and TLR-4 may be particularly important, as described in Supplemental Table 2. Given the current status of knowledge, the IO-specific DDPP gene set focuses on PD-L1, PD-L2, PD-1, CTLA-4, CD80, CD86, LAG-3, and TLR-4, together with specific markers of the presence of effector tumor-infiltrating immune cells: CD8A (cytotoxic lymphocytes T), CD16 (natural killer cells), and FOXP3 (T-reg cells). Many types of immune cells are involved in the activation and regulation of the immune system attack against tumor cells (APC, LyT, CD4+, etc.), but we focused on specific markers for infiltrating LyTc, NK, and T-reg that have the ability to recognize directly the tumor cells’ neoantigens coupled with major histocompatibility complex 1 (CMH1) and are directly targeting tumor cells.

A Pearson correlation analysis between differential gene expression of the selected genes, and the PFS was performed for the three patients treated with anti-PD-1 antibodies (Table 1) in the WINther trial. The example provided in Fig. 3a–c shows the DDPP intensity plots of three patients treated with IO (pembrolizumab (anti-PD-1), atezolizumab (anti-PDL-1), and nivolumab (anti-PD-1)); two patients had colon cancer (CRC) and one HN. Their clinical outcomes were ID 183 (CRC) PFS 61 months under treatment with pembrolizumab, ID 294 (HN) PFS 1.7 months under treatment with nivolumab in the WINTHER trial. The example provided in Fig. 3a shows the DDPP intensity plots of three patients treated with IO (pembrolizumab (anti-PD-1), atezolizumab (anti-PDL-1), and nivolumab (anti-PD-1)); two patients had colon cancer (CRC) and one HN. Their clinical outcomes were ID 183 (CRC) PFS 61 months under treatment with pembrolizumab, ID 294 (HN) PFS 1.7 months, and ID 270
predicted PFS under anti-PD-1 treatment was independent of the number of prior lines of therapy, if they were predicted PFS of the 82 patients from the WINTHER trial (for whom at the same value as the one observed for the predictor (PFS was 0.356%. The PFS of patients under afatinib treatment (n
six genes at each reiteration. Setting the threshold of significance was 0.04), the percentage of random significance with PFS was 0.356%.

Based on the six genes identified, we assessed “in silico” the predicted PFS of the 82 patients from the WINTHER trial (for whom no information was missing), agnostic of tumor type and independent of the number of prior lines of therapy, if they were treated with anti-PD-1 therapies. For 57 patients (59.5%), the predicted PFS under anti-PD-1 treatment was ≤6 months (with a majority <3 months); for 25 patients (30.5%) the predicted PFS under anti-PD-1 treatment was ≥6 months (of which 16 (19.5%) had predicted PFS > 24 months).

**DISCUSSION**

The expanded targeted and immune-oncology therapeutic landscape has revolutionized cancer care, as well as clinical drug development and requires the development of new combinatorial biomarker strategies to overcome the complexity of cancer, improve patients’ stratification and better individualize treatment selection.

Indeed, the investigation of networks of genes would contribute to improving the selection of drugs targeting major hubs that may collapse a network of dysregulated genes. Complication diagnostic (CDx) tests currently used in oncology practice to select therapies (e.g., used to detect specific DNA aberrations), have three main limitations: (1) CDx tests usually investigate single genes, outside of the context of pathways; for example, sensitizing mutations of EGFR provide an indication for the use of EGFR inhibitors, such as erlotinib, however the presence of a sensitizing mutation does not necessarily predict response to the treatment; (2) the increasing number of drug targets requires testing of multiple genes that quickly becomes unfeasible given the small amount of tissue typically obtained by needle biopsies; and (3) they provide a binary prediction whether patients might respond or not to a specific therapeutic regimen.

In an attempt to overcome the limitations of current oncology biomarkers, our transcriptomic approach allows (1) the exploration of the whole transcriptome (~20,000 genes), providing insight about the status of activation of almost all drug targets in the context of the network of genes or pathways; (2) the data can be obtained from a single assessment requiring very small amount of tumor and analogous normal tissues; and (3) the prediction of the duration of the PFS under a specific therapeutic regimen, as a continuous variable in contrast with binary predictions.

One of the main challenges of finding such new biomarkers is the small size of the available datasets, often involving fewer than 10 patients treated with the same drug, for whom both molecular profiles (from tumor and analogous normal tissues) and PFS data are available.

The work presented here overcomes the above mentioned limitations of current biomarker strategies and identifies the genes and generates linear regression that correlate with clinical outcome. In particular, the specific contribution of the DDPP and the features that distinguish it from other existing methods such as MLR or multiple Cox regression (MCR) consists in the use of nonparametric combination of the relative expression of multiple genes. Indeed, MLR and MCR investigate in the same model not only expression data, but also apply simultaneously coefficients to better differentiate genes impacting outcomes, compared to those that do not. The learning process defining such coefficients is a major limiting factor to these methodologies when using small datasets. DDPP on the other hand, in order to overcome this limitation empirically decides (1) how many features (genes) to include in the predictors rather than stopping when the fit is no longer improved, and (2) which basic parameter-free summation method best predicts the outcome. The final result of the DDPP model consists in defining the equations that link gene expression features to clinical outcome.

However, it is important to emphasize the limitation of the linear regression equations presented here, which are based on a prototype version of the DDPP. Indeed, the DDPP tools presented have the purpose of illustrating the concept and will require validation in a larger prospectively obtained dataset to confirm whether its empirical approach and applicability in any cohort size can be confirmed. Prospective studies will indeed assess the
ability of DDPP algorithm to test multiple summation methods in cohorts of different sizes.

One of the pillars, and potential limitations of the DDPP methodology is the investigation of tumor and analogous organ-matched normal tissue biopsies from the same patient, which is not the current practice in standard of care or in translational research and clinical trials. This raises the legitimate question of what is the likelihood of these findings to apply to routine oncology practice. This will be possible only if the DDPP predictor will demonstrate superiority to existing/current biomarkers models and provide a direct benefit for patients.

Use of matched tumor and normal biopsies is essential for accurate interpretation of the transcriptomic data, as it discards the transcriptomic genetic variability background noise in each patient, and lowers significantly the variance of transcriptomic measurements. We were able to use such transcriptomic and PFS data from the WINTHER trial database. The WINTHER trial remains the only clinical trial that used transcriptomics in a prospective clinical setting in addition to conventional DNA sequencing to help inform the treatment decision for patients with advanced cancer. WINTHER is also the first and only trial that used the dual biopsy strategy and proved its feasibility and safety, investigating both tumor and analogous normal tissue from the same patient, across a variety of solid tumors. The importance of obtaining good quality tumor and normal tissue biopsies enabling accurate transcriptomic investigations should be noted. All the biopsies used in this study passed stringent histology and RNA quality controls.

With the small cohort available, we explored the DDPP to assess correlations with PFS associated with the drugs everolimus, axitinib, trametinib, afatinib, experimental FGFR inhibitors, and anti-PD-1/PD-L1 therapies received by patients in the WINTHER trial. Remarkably, for all drugs tested, DDPP enabled identification of significant correlations between the differential expression of subsets of key genes and the PFS for each drug investigated. Our preliminary observations show that the DDPP biomarkers seem to be specific to the therapeutic regimens. It should be noted that the DDPP was agnostic of tumor type and independent of the number of prior lines of therapy and could also provide important insight in better understanding the clinical outcomes by identifying the genes with the highest contributing weight driving the correlations. Random testing performed for all drugs and in particular the data obtained from the leave-one-out experiments performed for everolimus, suggest that DDPP predictions are not likely to be statistical artefacts. Moreover, our data suggest that the subsets of genes selected, and the correlations obtained by their combined differential tumor versus normal tissues expression (vector summation) with the PFS for each therapeutic regimen may be specific for each drug and have a predictive value rather than a prognostic value, although this requires confirmation in larger studies.

Taken together, the data suggest the possibility that using a larger number of patients will allow us to generate a validated tool that may be useful to estimate with accuracy the PFS in a prospective clinical setting. Indeed, many drugs investigated in this report, have a narrow spectrum of approved clinical uses, given the prevalence of their pathways in tumor growth and spread, and the reason is probably related to the lack of reliable biomarkers to select patients who might have a therapeutic benefit.

To our knowledge potential biomarkers based on transcriptomics do not exist for the clinical use of everolimus or axitinib. Indeed, DDPP may provide a methodology and tools that would enable prediction of PFS for any drug (IO, or non-IO targeted therapeutics) or tumor type and in any therapy line. DDPP predictors could be used (pending further validation) to identify the patients who could have clinical benefit from the treatment with everolimus and axitinib that was not predicted by genomic alterations in the WINTHER trial.

The DDPP concept and methodology was tested also on other drugs: trametinib, afatinib, and two experimental FGFR inhibitors (BGJ398 and TAS-120), with a similar mechanism of action. However, the number of patients treated with each of these drugs (n=3), limits our ability to draw firm conclusions and the results are presented only to exemplify that DDPP concept and methodology may apply to any drug, and that similar trends were obtained as in the everolimus and axitinib examples.

We investigated a small cohort (n=3) of patients that received anti-PD-L1 therapies. Preliminary observations suggest that the main factors explaining differences in PFS under anti-PD-1 therapy are the degree of activation of TLR-4, and the balance between PDL-1, PDL-2, and CTLA-4 activation of the negative immune-blockade, together with the level of infiltration of the tumor by natural killer cells. Both DDPP intensity plots and vectorial summation correlations analyses identified TLR-4 as the most contributive gene to explain variations in PFS. Our preliminary observation suggests that the current panel of biomarkers used in clinical practice (tumor mutation burden, microsatellite instability, and PDL-1 status) could be complemented with other potential biomarkers, such as TLR-4. Persistent activation of TLR-4-induced inflammatory signaling in chronic inflammatory conditions can also contribute to carcinogenesis. Experimental evidence suggests association of anti-TLR-4 with anti-PD-L1 treatments could be of interest with the aim to increase the fraction of patients who could benefit from IO treatments.

Validation of DDPP will require further prospective investigation, ideally using biospecimens and clinical data obtained from patients participating in prospective clinical trials, in which implementation of dual tumor and normal biopsy and integration of transcriptomics are feasible.

In conclusion, the clinical records available from the WINTHER trial, and the unique transcriptomic dataset obtained from tumor and organ-matched normal tissue biopsies were essential to enable correlations with clinical outcome under treatment, with TKI inhibitors and IO. The DDPP is potentially a novel global biomarker model that can apply to any type of drug (IO or non-IO targeted drugs) alone or in combination, agnostic of tumor type, and can lead, pending further prospective validation, to a new approach to optimal treatment selection for patients with cancer.

METHODS

Dataset

The DDPP prototype was developed exploiting the transcriptomics data from the WINTHER trial (NCT 01856296—approved in France (ANSM), Spain (Agencia Española de Medicamentos y productos sanitarios), Israel (Ministry of Health), Canada (Health Canada), and USA (FDA)). The full methodology of transcriptomic assessment and patient treatment is described in the WINTHER trial published in Nature Medicine. Detailed clinical and biological information for each patient selected for this study is available in Table 1.

Patients who had provided written consent underwent a dual biopsy from the metastasis and from the historically matched normal tissue. The interventional radiologist selected for biopsy one of the non-necrotic and non-calcified metastatic lesions. The image-guided biopsies were performed under standard operating rules and guidelines using adequate devices, such as trucut 18 gauges needles placed at the periphery of the lesion biopsied. For example, in the case of a liver metastasis biopsy from a rectal adenocarcinoma, the matched normal tissue was the rectal mucosa obtained separately by rectal endoscopy. All biopsies obtained were stored in cryogenic tubes containing RNALater, a stabilizing reagent preventing degradation of nucleic acids (without the need for freezing in the imaging facilities), and preserving structural morphology to enable pathology review. Biopsies were embedded in OCT wax. Slices of 0.5 microns were cut and stained with hematoxylin and eosin, in order to assess the percentage of tumor cells and identify any contamination with adjacent normal tissues.18

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tissues. When necessary, microdissection was performed to enrich tumor content to the required threshold (≥50% of tumor content).

Tissue biopsies that had passed quality control were then lysed by a polytron, homogenized in a lysis buffer RLT Plus provided in the kit, and DNA was obtained with a specific affinity silica matrix column, specifically retaining the DNA, whereas RNAs and proteins were collected from the through flow. DNA was washed and eluted. The through flow containing RNAs was mixed with tri-reagent, and subsequently RNA was obtained by isopropanol precipitation. This procedure enables collection of all types of RNAs, including messenger RNAs and small microRNAs species. RNAs pelleted through centrifugation were washed with ethanol 75%, and solved in nucleos-free water. Quality control for DNAs and RNAs was performed using spectrophotometry absorbance (NanoDrop) and through electrophoresis, using lab-on-a-chip technology from Agilent Technologies.

Gene expression direct comparison of tumor and normal tissue RNAs was performed using Agilent inkjet printing 8x 60k oligo-arrays and dual color technology, using standard operating procedures and reagents supplied by Agilent Technologies. A total of 100 ng of each tumor and normal tissue total RNA was used to generate double-stranded complementary DNA, using MMLV Reverse transcriptase and oligo DT primers coupled with the promoter sequence of T7 RNA. Probe labeling and linear amplification were generated using Agilent Technologies reagents and T7 RNA pol that generated labeled complementary labeled RNAs (tumor labeled with Cy5 and normal with Cy3RNAs with Cy5). After fragmentation and purification of labeled RNA, hybridizations were performed as dual dye swapped (direct and inverted labeling) experiments with direct co-hybridization of equal amounts of labeled tumor and normal probes. After washing drying, microarrays were read using Agilent 2000 scanner version C. After processing with Agilent Feature extraction software, data were used for direct comparison of intensities and editing of a report containing a data quality information, identity of RNAs differentially expressed (overexpressed or underexpressed in tumor versus normal, and providing for each gene fold changes and intensities together with p values for each measure, type of structural abnormalities (amplifications/deletions), threshold, fold changes, and intensities).

DDPP algorithm

The concept of DDPP algorithm is derived from the Euclidian linear hyperspace model, in which the distance between different outcomes can be defined using multiple vectors (or coordinates). The cornerstone of the Euclidian model’s application to precision oncology and to DDPP is the identification and summation of the optimal coordinates, which are defined as the mechanism-based key genes that govern sensitivity to each of the targeted medications investigated. The DDPP methodology that applies to any type of drugs is based on the following steps:

1. **Identification from the literature**, of the key genes governing the sensitivity for a given therapeutic regimen and exploring their differential expression in tumor compared to analogous normal tissue. This dramatically reduces the number of features (genes) that are considered in subsequent steps. As we observed only six instances (patients with known PFS values under treatment with everolimus), this is a crucial stage: without dramatic decrease in the number of genes, overfitting (i.e., the development of a perfect model that is specifically tailored to the given dataset used for training), is almost guaranteed. Identification of key genes involved in drug’s mechanisms of action, based on recent literature and based on the FDA US Prescribing Information (USPI).

a. **Everolimus**—key genes: PIK3CA, PIK3CB, AKT1, MTOR, FKB11A, RPS6KB1, EIF4EBP1, HIF1A, TSC1, TSC2, AKT2, RPTOR, PTEF, RHEB, MLST8, NICTOR, and VEGFA.

b. **Axitinib**—key genes: VEGFA, VEGFB, VEGFC, PDGFA, PDGFB, FLT1 (VEGFR1), KDR (VEGFR2), FLT4 (VEGFR3), PDGFRα, PDGFRβ, KIT, FLT1 (VEGFR1), and FIGF.

c. **Trametinib**—key genes: MEK1 (MAP2K1), MEK2 (MAP2K2), ARAF, BRAF, RAF1, ERK (MAPK3), ERK2 (MAPK3), MAPK10, KRAS, HRAS, NRAS, KSR1, and RAP1A.

d. **Afatinib**—key genes: EGFR, ERBB2, ERBB3, ERBB4 and their ligands EGF, TGFα, AREG, EREG, HBBEGF, BTC, NRG1, NRG2, NRG4.

e. **FGFR inhibitors**: FGFR1, FGFR2, FGFR3, FGFR4 and the FGFR ligands 1, 2, 3, 4, 5, etc.

f. **Anti-PD-1/PD-L1**—key genes: PDL-1, PDL-2, PD-1, CTLA-4, CD28, CD80, CD86, LAG-3, and TLR-4, together with specific markers of the presence of effector tumor-infiltrating immune cells: CD8A (cytotoxic lymphocytes T), CD16 (natural killer cells), and FOXP3 (T-reg cells).

2. **Selection of the patients** with available transcriptomics data and clinical outcome (PFS) under treatment with each drug available. Minimum three patients are required. Everolimus (n = 6); axitinib (n = 5);

a. Preferably, patients with non-censored PFS were selected for investigations.

b. Only two patients had censored PFS under treatment (ID 203 treated with everolimus and ID 183 treated with pembrolizumab who had exceptional PFS >60 months, and were still under treatment). We did not discard them from the analysis, but considered them de-censored.

3. **Comparing (empirically) the performance of all top-ranking subsets**: our method is based on ranking each of the selected features (genes), based on their relative association with the dependent variable (i.e., PFS). This step can be performed in multiple ways: genes can be ranked by the strength of their univariate linear regression with PFS (e.g., Pearson) or univariate regression model that is specifically tailored to the given dataset used for training), is almost guaranteed. Identification of key genes involved in drug’s mechanisms of action, based on recent literature and based on the FDA US Prescribing Information (USPI).
investigation, both methods are too likely to over fit the model to the data. As both rely on parametric linear combination for combining the features into a single method.

We propose the Digital Display Precision Predictor (DDPP) algorithm, as an alternative way for features summation that is parameter free. DDPP is different from these approaches as it empirically chooses the summation method that gives the best model, considering in the current version five basic, parameter-free summation methods: sum, mean, product, absolute product, and scalar median. It differs from the methods described above in not trying to estimate any coefficients in the summation step, making it especially suitable for small datasets, as it is less likely to over fit.

In conclusion, DDPP is unique in two levels:

- It empirically decides how many features (genes) to include in the predictors rather than stopping when the fit is no longer improved.
- It empirically decides which basic parameter-free summation method best predicts the outcome.

5. Combined gene expression (parameter-free feature summation): in a previous publication, we explain how we derive $F_i$ from two color arrays. Briefly Equation (1) describe how feature $i$ is calculated for gene $g$:

$$F_{ig} = \frac{1}{N_g} \sum_{x} \log_{1.1} f_{gx} \cdot \log_{1.1} I_{g,tumor}$$

where $F_{ig}$ is used as the measurement for gene $g$ in individual $i$, $T_{ig}$ and $N_g$ are the measured intensity of gene $g$ in individual $i$ correspondingly. In order to get a single value from multiple features (genes), we combined for each individual all genes values ($F_{ig}$, $f_{ig}$, $I_{g,tumor}$, $I_{g,normal}$) using different approaches: (1) mean: $\overline{F}_i$, (2) sum: $\sum_{x} F_{ig} / (3)$ median; (4) fold: $\frac{I_{g,tumor}}{I_{g,normal}}$, (5) fold_abs: $|\frac{I_{g,tumor}}{I_{g,normal}}|$.

6. To define the optimal “n” genes investigated we interrogated the correlation between gene expression and the PFS. For each drug, a Pearson correlation test was performed between $F_p$, the fold change multiplied by the intensity of (tumor and normal) of a single gene $g$ (gene from the list of key genes of the drug) with the PFS for all the patients treated with the drug.

$$F_p = \log_2(\text{fold change tumor vs. normal}) \times \log_{1.1} (\text{Intensity of tumor})$$

The $F_p$ with the most significant correlated gene (by $p$ value and correlation coefficient $r$) was driven to decide whether to continue with fold change multiplied by the intensity of the tumor or fold change multiplied by the intensity of the normal matched tissue. The key genes were then ranked based on the Pearson $p$ value and correlation coefficient $r$ such that the gene with the highest correlation between $F_p$ and the PFS was ranked first. Then, we added single genes by the following manner: the second most ranked gene was added to the first most ranked and the $F_{ig,2}$. This resulted with vector space which contains two vectors. Similarly, the third most ranked gene was added to the second highest ranked genes, and resulted with vector space with three vectors. The vector summation continued in a similar manner until we obtain $n$ different combinations of genes, while $n$ is the number of genes in the set and each combination denotes a vector space with $n$ vectors in it. In order to get a single value for each vector space, we tested the five different basic mathematical operations on the vectors in each vector space mentioned above.

Then, a Pearson correlation test was performed between the calculated transcriptomic value for each vector space with the PFS of the patients treated with the drug. The Pearson correlation results were ranked again by $p$ value and correlation coefficient $r$. The number of genes in the set which was the most correlated with the PFS was indicated as the optimal “n” coordinates. In order to assess the likelihood of getting a significant correlator by “n” genes, we run an analysis with 100k random “n” genes and tested how the transcriptomic value derived from a vector space which includes $F_{ig,1-n}$, of these genes were correlated with the PFS. Significant results were considered by a threshold of absolute $R$ value of 0.9 or above, and $p$ value of 0.05 and below.

Applying any of the mathematical operations listed above resulted with a single value that combines the transcriptomic expression of all targetable genes for each patient. These values were used in order to generate the linear regression model with the PFS observed.

7. Selecting the best correlators with PFS for each drug and computing a linear regression model to transform the best correlator into a predictor for a single drug. Supplemental Table 3 shows the log2 fold change (intensity in tumor versus intensity in normal) multiplied by the intensity of tumor of selected eight genes for patients treated with everolimus. The expression values were used to calculate DDPP values based on fold_abs, which is absolute value of the multiplication of all the gene’s values in each patient. The DDPP values are shown in Supplemental Table 4. Then, we used the DDPP values and the real PFS values as shown in Supplemental Table 5 for Pearson correlation with 95% confidence interval analysis and linear regression model analysis (which provided the linear equation). The results of these analyzes are shown in Supplemental Fig. 3.

8. Linear regression equations developed to link gene expression with clinical outcome under specific treatments:

- Everolimus—Equation 1: $Y = 1.499e-13X + 3.134$, where $X$ is the absolute value of the fold of log2(fold change tumor versus normal) multiplied by log1.1(Intensity_Tumor) of each value for each of the eight genes (AKT2, TSC1, FKB-12, TSC2, RPTOR, RHEB, PIK3CA, and PIK3CB) and $Y$ = PFS in months.
- Axitinib—Equation 2: $Y = 2.014e-02X + 4.36$, where $X$ is the sum of log2(fold change tumor versus normal) multiplied by log1.1(Intensity_Normal) of each value for each of the two genes (KIT, KITLG), and $Y$ = PFS in months.
- Trametinib—Equation 3: $Y = -6.872e-15X + 7.745$, where $X$ is the fold of log2(fold change tumor versus normal) multiplied by log1.1(Intensity_Tumor) of each values for each of the nine genes (ERK2, ARAF, CRAF, MEK1, MEK2, HRAS, ERK1, MAPK10, and KRAS), and $Y$ = PFS in months.
- Aftinib—Equation 4: $Y = -4.558e-02X + 2.549$, where $X$ is the sum of log2(fold change tumor versus normal) multiplied by log1.1(Intensity_Tumor) of each value for each of the two genes (NRG4 and NRG2), and $Y$ = PFS in months.
- FGFR inhibitors—Equation 5: $Y = 5.273e-02X + 5.135$, where $X$ is the sum of log2(fold change tumor versus normal) multiplied by log1.1(Intensity_Normal) of each value for each of the five genes (FGF10, FGF16, FGF5, FGF2, and FGF13), and $Y$ = PFS in months.
- Anti-PD-1—Equation 6: $Y = 7.856e-10X + 1.583$, where $X$ is the value of the fold of log2(fold change tumor versus normal) multiplied by log1.1(Intensity_Normal) of each value for each of the six genes (TR-L4, PDL-2, PDL-1, CD16 (NK), CTLA-4, and CD28), and $Y$ = PFS in months.

Reporting summary
Further information on research design is available in the Nature Research Reporting Summary linked to this article.

DATA AVAILABILITY
Expanded clinical and biological information for each patient is available in the data file Patient_demographic_clinical_data.csv which expands on the information presented in Table 1. The data file DPP_WINTHER_Expression_Data.csv supports Figs. 1–3, Supplementary Tables 1, 3–5, and Supplementary Figs. 1–3. The transcriptomic expression data are available from Gene Expression Omnibus: https://identifiers.org/geo:GSE168621 (ref. 30). The data generated and analyzed during this study are described in the following data record: https://doi.org/10.6084/m9.6.figshare.14166731 (ref. 29).

CODE AVAILABILITY
Codes are available upon request to the corresponding author.

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AUTHOR CONTRIBUTIONS

V.L., S.M., N.G., A.S., J.F.M., G.M., I.B.N., M.S., A.P., A.M.T., E.R., R.K. and R.S. contributed with conception of the presented idea and design of the project. R.B., A.O., F.W., I.B., J.T., E.F., G.B., C.K., B.S. and J.C.S. contributed to the implementation of the project with patient follow up and clinical outcome database update. V.L., S.M., A.S. and E.R. performed the bioinformatics computations and statistical analysis. V.L., S.M., N.G., A.S., M.S., A.P., J.R., B.S., A.M.T., R.K. and R.S. contributed to the interpretation of the results. V.L., S.M., C.B., A.M.T., J.F.M., R.K. and R.S. took the lead in writing the manuscript. All authors provided critical feedback and helped shape the research, analysis and manuscript, and approved the final version.

COMPETING INTERESTS

Dr. Vladimir Lazar, Catherine Bresson, and Fanny Wunder, are full-time employees of Worldwide Innovative Network (WIN) Association—WIN Consortium. Worldwide Innovative Network (WIN) Association—WIN Consortium is the owner of the patent family entitled Digital Display. The inventors are Dr. Vladimir Lazar, and Shai Magidi. Shai Magidi receives consultancy from Worldwide Innovative Network (WIN) Association—WIN Consortium. Dr. Jean-François Martinis is a full-time employee and stockholder of Pfizer Inc. Dr. Amir Onn receives consulting fees from Roche Israel, MSD Israel, Boehringer Ingelheim, and AstraZeneca. Dr. Giorgio Massimini is employee and shareholder of Merck KGaA, consulting for The Hellday Foundation (for free), shareholder of Hoffmann-LaRoche and Calithera. Irene Braha receives consulting fees from Orion Pharma, receiving speaking fees from BMS, AstraZeneca, and Merck Serono, and is principal investigator and receives funding for clinical trials from AstraZeneca, BMS, Celgene, Gilkin, GSK, Janssen, KURA, MSD, Novartis, Orion Pharma, and Pfizer. Dr. Raanan Berger receives consulting fees from Roche, MSD, AstraZeneca, and BMS. Dr. Josep Tabernero declares scientific consultancy role for Array Biopharma, AstraZeneca, Bayer, BeiGene, Boehringer Ingelheim, Chugai, Genentech, Inc., Genmab A/S, Halozyme, Imugene Limited, Infiltration Biosciences Limited, Ipsen, Kura Oncology, Lilly, MSD, Menarini, Merck Serono, Merrimack, Merus, Molecular Partners, Novartis, Peptomyc, Pfizer, Pharmacies, ProteoDesign SL, Rafael Pharmaceuticals, F. Hoffmann-LaRoche Ltd., Sanofi, SeaGen, Seattle Genetics, Servier, Symphogen, Taiho, VCN Biosciences, Biocartis, Foundation Medicine, HaloDX SAS, and Roche Diagnostics. Dr. Enriqueita Felip receives advisory board and/or speaker’s bureau fee from AbbVie, Amgen, AstraZeneca, Bayer, BluePrint Medicines, Boehringer Ingelheim, Bristol-Myers Squibb, Eli Lilly, GSK, Janssen, Mediscape, Merck KGaA, Merck Sharp & Dohme, Novartis, Pfizer, pmxME Oncology, Puma Biotechnology, Sanofi, Genzyme, Springer, Takeda, and Touchimove: on the board of Grifols, Independent member. Receives research funding from Fundacion Merck Salud, Grant for Oncology Innovation (GOI) EMD Serono. Prof. Angel Porgador declares scientific consultancy role for PINK Biopharma. Dr. Gerald Batist collaborates in formal clinical trial contracts, IITs and in joint grants funded by Canadian and Quebec governments with Roche, Merck, Novartis, AstraZeneca, Bayer, Esperas, Aurora, Caprion, and MRM. Dr. Benjamin Solomon received honorarium for Advisory Board for AstraZeneca. Prof. Apostola Maria Tsimberidou receives research funding (her institution) from Immatics Biotechnologies, Parker Institute for Cancer Immunotherapy, OBI Pharmaceuticals, Karus Therapeutics, Ltd., Tempus, Inc. CPRT, Tvardi Therapeutics, EMD Serono, and Boston Biomedical. Inc. She also receives support from Philanthropic Funds. She has received consulting fees from Genentech, Covance, and Tempus (past 2 years). Dr. Razelle Kurzrock receives research funding from Genentech, Merck Serono, Pfizer, Boehringer Ingelheim, TopAlliance, Takeda, Incyte, Debiapharm, Medimmune, Sequenom, Foundation Medicine, Konica Minolta, Grifols, Omnisecq, and Guardant, as well as consultant and/or speaker fees and/or advisory board for X-Biotech, Neomed, Pfizer, Actuate Therapeutics, Roche, Turning Point Therapeutics, TDIQ, Oncobio, and Bica Therapeutics, Inc., has an equity interest in IdbyDNA and CureMatch Inc, serves on the Board of CureMatch and CureMetrix, and is a co-founder of CureMatch. Dr. Richard L. Schilsky ’s institution (ASC0) receives research grants in support of a clinical trial that he directs from the following companies: AstraZeneca, Bayer, Boehringer Ingelheim, Bristol-Myers Squibb, Genentech, Lilly, Merck, and Pfizer. The remaining authors declare no competing interests.

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