TP53 mutations and number of alterations correlate with maximum standardized uptake value (SUVmax) determined by positron emission tomography/computed tomography (PET/CT) \([^{18}\text{F}]\) fluorodeoxyglucose (\(^{18}\text{F}-\text{FDG}\) PET)

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ABSTRACT

Background: Our study explored the relationship between the molecular changes in cancer and the maximum standardized uptake value (SUVmax) determined by positron emission tomography/computed tomography (PET/CT) with \([^{18}\text{F}]\) fluorodeoxyglucose (\(^{18}\text{F}-\text{FDG}\)).

Results: A higher SUVmax correlated with TP53 alterations, but not with histologic diagnosis or other gene/pathway mutations or copy number alterations. In data from breast, lung and colon cancer, patients with the highest SUVmax show more genomic anomalies compared to those with the lowest SUVmax \((P < 0.005)\).

Conclusions: A higher SUVmax on \(^{18}\text{F}-\text{FDG}\) PET/CT is associated with TP53 tumor suppressor gene anomalies and the presence of more genomic anomalies. Since TP53 alterations and high SUVmax both correlate with a poor prognosis, the underlying mechanism/implications of this association merit further study.

Methods: Overall, 176 patients with diverse cancers had a tumor biopsy within 6 months after a PET/CT image for SUVmax measurement. The biopsy was interrogated by next generation sequencing (182 to 315 genes). TP53, EGFR, ALK, MYC, MET and FGF/FGFR genes and DNA repair, PI3K/Akt/mTOR (PAM), MEK, CYCLIN, and WNT pathway genes were analyzed.

INTRODUCTION

Cancer is the second most common cause of death in the US, exceeded only by heart disease \([1]\). Current trends demonstrate a decline in four of the most common cancer types—lung, colorectal, breast and prostate \([1–3]\). Such improvements reflect both the earlier diagnosis of certain cancers and improvements in treatments.

One area of intense research is the use of comprehensive genomic profiling with the use of next generation sequencing \([4–5]\). The goal is to use this information to expand treatment options by matching an individual patient with targeted therapies and clinical trials that are relevant to the molecular changes in the tumor \([6–7]\).
RESULTS

Patients

We studied 176 patients (109 women, 67 men) aged 23 to 85 years (mean age = 57 years) with the diagnosis of: breast cancer 41 (23%), lung cancer 44 (25%); gastrointestinal cancer 39 (22%), and a variety of other tumor types. Due to the retrospective nature of this study, the patients are in different stages of their cancers and the lesion biopsied included primary and metastatic lesions.

SUVmax as a linear variable

SUVmax as a linear variable was analyzed using a univariable test—the Mann-Whitney test. The overall mean for SUVmax was 8.30 ± 7.65. The overall median for SUVmax in the N = 176 patients = 6.4 (5.8–7.7). There was no significant difference in the SUVmax in the patients when grouped by their diagnosis: breast cancer 7.23 ± 4.12, lung cancer 7.97 ± 5.09, and GI cancers 7.40 ± 4.54.

Genomic anomalies (Table 1)

TP53 Gene

Eighty-four of the 176 patients (48%) had alterations in TP53 and a significantly higher median SUVmax of 7.85 compared to unaltered TP53 with a reference median SUVmax of 5.85 (P = 0.023).

DNA repair genes (BRCA, BRIP, ATM, MMR, MSH, MLH)

Twelve of 176 patients (7%) had alterations in DNA repair genes. These 12 patients demonstrated a median SUVmax of 9.65 compared to the median SUVmax of 6.15 in the 164 patients with other gene alterations (P = 0.130) (but the affected number of patients is small).

EGFR gene

Thirty-six of 176 patients (20%) had EGFR gene anomalies. The median SUVmax of these 36 patient was 6.5 which was not significantly different than those without EGFR gene anomalies (P = 1.0).

PI3K/AKT/MTOR (PAM) genes (PTEN, PIK3CA, AKT, TSC, CCNB1, MTOR, FBXW2, NF2)

Fifty-eight of the 176 patients (33%) had abnormalities in PAM pathway genes. The median SUVmax for these 58 patients was 7.75 compared to 6.05 in the patients without abnormal PAM pathway genes. This was not statistically significant (P = 0.062).

MEK genes (RAS, RAF, MAPK, CNAS)

Thirty of the 176 patients (17%) had MEK gene anomalies. Their median SUVmax was 7.75 as compared to 6.20 without the MEK gene anomaly. This was also not significantly different (P = 0.291).

CYCLIN genes (CCND, CDK, CDKN, RB)

Sixty-six of the 176 patients (38%) had abnormalities in CYCLIN pathway genes and a median SUVmax of 6.15 compared to 6.55 in the patient without a CYCLIN gene abnormality. These means were not significantly different (P = 0.483).

WNT genes (APC, CTNNB, NOTCH)

Twenty-five of the 176 patients (14%) had abnormalities in WNT pathway genes and a corresponding median SUVmax of 6.0 compared to a median SUVmax of 6.5 in patients without WNT pathway abnormalities. These median SUVmaxes were not statistically significant, (P = 0.842).

MYC gene

Twenty-five of the 176 patients (14%) had MYC gene anomalies and a corresponding median SUVmax of 7.9 as compared to the 151 patients without MYC gene anomalies with a median SUVmax of 6.2. This was not significantly different (P = 0.431).

FGF/FGFR genes

Twenty-eight of the 176 patients (16%) had FGF/FGFR gene anomalies and a median SUVmax of 5.35 as compared to the 148 patient without anomalies and an median SUVmax of 6.8. This was not significant (P = 0.380).

ALK and MET gene

Three of the 176 patients (2%) had ALK gene anomalies. Four of the 176 patients (2%) had MET gene anomalies. Only variables with ≥10 patients were included and, thus, these gene alterations were not assessed in the univariable analysis.

Number of alterations

The average of the three highest and three lowest SUVmaxes and the presence of the various genomic anomalies in the three patient groups—breast cancer, lung cancer, and colon cancer are shown in Table 2. In the three cancers, the three patients with the highest SUVmax show more genomic anomalies compared to the three patients with the lowest SUVmax (P < 0.005).
DISCUSSION

Oncologic therapies are evolving with increasing emphasis on the evaluation of multiple genomic alterations within the biological pathways driving tumorgenesis, with the idea of providing molecularly targeted therapy. $^{18}$F-FDG PET/CT is widely accepted as a standard of care for tumor staging/restaging as well as the modality of choice to detect malignancy and predict prognosis. In addition, $^{18}$F-FDG PET/CT often determines and guides therapeutic decisions and also serves as a way to monitor response to therapy.

The purpose of our study was to determine whether or not $^{18}$F-FDG PET/CT SUVmax correlated with specific...
CONCLUSIONS

A higher SUVmax on $^{18}$F-FDG PET was associated with oncogenic alterations in the $TP53$ tumor suppressor gene. Abnormalities in $TP53$ and increased SUVmax are each considered poor prognostic factors in patients with cancer. A higher SUVmax also correlated with the presence of more oncogenic alterations. Further investigation of the mechanism underlying correlations between genomic anomalies and PET imaging results is warranted.

MATERIALS AND METHODS

Patients

We studied 176 consecutive patients with available data who underwent $^{18}$F-FDG PET/CT six months or less before a biopsy for genomic profiling. The SUVmax of the biopsied lesion was obtained from the PET/CT imaging. Molecular testing data was analyzed to determine the presence of common DNA alterations. For our study, we included $TP53$, DNA repair genes ($BRCA$, $BRIP$, ATM MMR, MSH, MLH), $EGFR$, $PI3K/Akt/mTOR$ (PAM) pathway genes ($PTEN$, $PIK3CA$, AKT, TSC, CCNB1 MTOR, FBXW2), $CYCLIN$ (CCND, CDK, CDKN, RB), WNT pathway ($APC$, CTNNB, NOTCH), $ALK$, $MYC$, and FGF/FGFR genes. This study was performed and patients consented in accordance with the guidelines of the UCSD Internal Review Board (PREDICT [Profile Related Evidence Determining Individualized Cancer Therapy], protocol; NCT02478931).

$^{18}$F-FDG PET/CT

A combined $^{18}$F-FDG PET/ CT scanner (General Electric Discovery VCT PET/CT, Waukesha, WI) was used to perform whole body imaging. Images were interpreted according to standard methods. Whole-body CT covers a region ranging from the head to the mid-thigh. After fasting for at least four hours, patients received an intravenous injection of $^{18}$F-FDG [10–20 mCi]. Blood glucose was checked in all patients before performing $^{18}$F-FDG PET/CT and no patient had a blood glucose level >160 mg/dl. About 50 minutes later, CT scanning was conducted and whole-body emission PET scanning was performed. Attenuation-corrected PET images were reconstructed with an iterative reconstruction algorithm. $^{18}$F-FDG PET/CT images were generated for review on a workstation.

Qualitative analysis of $^{18}$F-FDG PET/CT

Quantitative analysis was performed on the institution’s pictures archiving and communication system (PACS), (AGFA Impax 6.3, Mortsel Belgium). The PACS software was used to draw a single region of
interest (ROI) in the area of the lesion with most intense uptake to determine the SUVmax. The SUVmax was determined according to the standard formula, with activity in the region of interest (ROI) being calculated as mCi/ml divided by the injected dose and normalized to the patient's body weight. The SUVmax was defined as the maximum activity within the ROI.

Genomic analysis

Genomic analysis was performed using a clinical next generation sequencing (NGS) based assay (182 to 315 genes) (FoundationOne™, Foundation Medicine Inc., Cambridge, MA), which includes detection of base substitutions, insertions, deletions, copy number alterations, and selected gene fusions. We included EGFR, PI3K/Akt/mTOR (PAM) pathway genes, CYCLIN, WNT pathway, ALK, MYC, and FGF/FGFR genes.

Author contributions

Geraldine H Chang: collected and analyzed PET image data; Razelle Kurzrock: data analysis; Lisa Tran: collected and analyzed tumor markers; Maria Schwaederle: statistical analysis of data; Carl Hoh: assisted in interpretation of quantitative PET data.

CONFLICTS OF INTEREST

Dr. Kurzrock has research funding from Genentech, Merck Serono, Pfizer, Sequenom, Foundation Medicine, and Guardant Health, as well as consultant fees from X-Biotech and Actuate Therapeutics and has an ownership interest in Curematch, Inc.

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